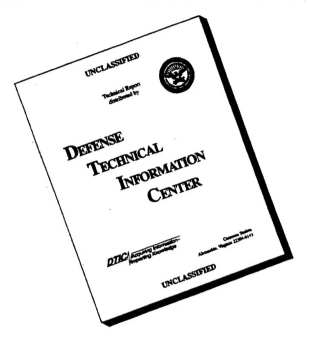
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## Adsorption and Catalysis on Carbonaceous Adsorbents - Electronic Factors

by

Edward George Marchand

#### A DISSERTATION

Submitted in partial fulfillment of the requirements

for the degree of

DOCTOR OF PHILOSOPHY

(Environmental Engineering)

MICHIGAN TECHNOLOGICAL UNIVERSITY

1996

This dissertation, "Adsorption and Catalysis on Carbonaceous Adsorbents - Electronic Factors," is hereby approved in partial fulfillment of the requirements for the degree of DOCTOR OF PHILOSOPHY in the field of Environmental Engineering

RROGRAM: Environmental Engineering

Dissertation Advisor: Dr. David W. Hand

Chair, Environmental Engineering Ph.D. Committee:

Dr. John C. Crittenden

#### **ABSTRACT**

Two carbonaceous adsorbents made from polymeric materials were used in adsorption and catalytic oxidation experiments in the lab and the field. One was a phenolaldehyde based (Novoloid) fiber, Kuractive, used in a cloth or fabric form. The other was a sulfonated polystyrene divinylbenzene cross-linked polymer adsorbent, Ambersorb® 572. Catalytic oxidation of trichloroethylene (TCE) at 110-250 °C and space velocities of 36,000 to 73 hr<sup>-1</sup> (Empty Bed Contact Time, EBCT, 0.1 - 49 s) over plain and transitionmetal-oxide impregnated Ambersorb 572 demonstrates the potential for this catalyst/support to convert TCE at low temperatures. The first step in the catalytic process appears to be the addition of a proton from the acid catalysis sites to the TCE, followed by non-specific attacks by the resulting carbenium ion or radical. High conversions were maintained at temperatures as low as 110 °C, although there were numerous by-products. At temperatures above 180 °C the plain adsorbent/catalyst was superior in performance over the metal-impregnated catalysts under the conditions in this study. Water in the feed stream has an immediate, reversible, poisoning effect on the performance of MTU-CAT, a thermally modified Ambersorb 572, and the transition metal-impregnated catalysts used in this study. However, the water poisoning effect was not noticed when the catalyst was the plain support, Ambersorb 572.

Resistive or Joule heating was used to heat both adsorbents. This thermal energy can reduce the local humidity from the incoming gas without heating the entire air stream, regenerate the adsorbent, or provide the activation energy for the oxidation of the desorbing organics on catalyst sites in the adsorbent. At low currents (tens of milliamps at 2-3 V D.C.) the thermal energy applied to the carbon cloth can alter the local relative humidity, favoring adsorption of trichloroethylene over water, increasing the adsorption capacity of the adsorbent by a factor of 4.5 times. However, initial cost estimates favor the use of preheating the air, instead of Joule heating the adsorbent bed, to reduce the relative humidity to maintain high carbon capacities.

Adsorption of organic compounds onto the carbon fibers changes the resistivity of the materials. Different compounds altered the resistance in different ways and at different rates. Methanol, a polar compound, rapidly changed the resistance, while toluene, a non-polar compound, was the slowest to alter the resistance. Trichloroethylene initially decreased the resistivity, and then increased it over time. The change in resistance upon adsorption was found to be concentration dependent. The polarizability of the compound appears to be the key to the rate of change in the resistance. In humidified streams competition for adsorption is evident from the resistance profiles over time. The results of this work lead to the development of an inexpensive analytical device for monitoring the performance of gas-phase adsorption systems.

During a limited study a dynamic electric field (0.5-2 Amps, or 0.4-1.3 W/g) was applied to a catalyst bed of Ambersorb 572 at 180-200 °C. It was found to alter the composition of the effluent of the catalytic oxidation reaction of trichloroethylene (TCE). It is unclear if electrooxidation (transfer of electrons) is responsible or the change is merely a side effect of charging the surface of the adsorbent catalyst under electrodynamic conditions which would impact the stability of the carbenium ion. Under a static electrical field (+1000 V DC) flow-through tests in a lab reactor with TCE and field tests using a contaminated stream from a soil vapor extraction system indicate direct electrooxidation of organics may be occurring.

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Finally (why do we save the most important for last?) I wish to thank God for His grace and goodness. With His help I have overcome major obstacles. All praise and honor for this work are to Him who has redeemed my soul.

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#### **PREFACE**

This dissertation was written as a series of papers to be published in appropriate professional journals. As such, Chapters 2, 3, and 4 are written to stand alone and contain duplicative information, especially about the materials and methods used in the experiments.

Appendix A contains the raw data from the nitrogen adsorption/desorption BET done by Porous Materials Inc., (Ithaca, NY). Appendix B contains all the Quattro Pro spreadsheets giving the data supporting the figures. Appendix C contains supplemental information on the catalytic oxidation over the metal oxide impregnated catalysts. The information is summarized in Chapter 2 to keep it to a reasonable length as a publishable paper.

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This dissertation was written using the following software tools: Harvard Graphics, Quattro Pro, and Word Perfect.

#### Chapter 1

#### Introduction

#### 1.1 BACKGROUND

Industrial and remediation processes generate large quantities of air contaminated with organics. The U.S. Environmental Protection Agency (EPA) estimated 1.22 billion pounds of organics were released from point and non-point sources to the air in 1993 from U.S. facilities (US EPA, 1993). Clean Air regulations promulgated by the EPA place limits on the types and amounts of organics that can be released to the environment. Control technologies, especially for dilute waste streams with chlorinated organics, can be expensive and become the major cost involved with the process. At a groundwater contamination site the fuel costs for the catalytic oxidation unit used to treat the air stripper off-gas was almost 80% of the total utilities cost of \$0.36/1000 gallons water treated (Hylton, 1992).

Technologies currently used to limit the release of the organics to the air include gas phase adsorption onto granular activated carbon (GAC) or combustion, either in a direct flame incinerator at high temperatures (>2000 °C) or over a catalyst at moderate temperatures (200-500 °C). If carbon is used it must periodically be replaced or regenerated, with the resulting stream requiring further treatment/disposal. In combustion systems higher temperatures and/or longer time in the reactor (residence time) allow for complete conversion of the chlorinated organics to water and mineral acids such as HCl. This complete destruction is referred to as mineralization. Partial conversion of the molecules can result in products that are more toxic than the parent compound (such as phosgene from the oxidation of trichloroethylene).

#### 1.2 TARGETED TOXIC CHEMICAL SOURCES

The sources of chemicals are almost as varied as the number of chemicals in existence. The sources targeted for this research work are from the equipment operation/maintenance processes and remediation efforts at a typical Air Force Base. Volatile organic compounds (VOCs) released from these operations require off-gas control. The compounds and the processes include 1) engine cleaning compounds from a cleaning vat, 2) paint booth vapors from an exhaust unit, or 3) chlorinated and non-chlorinated VOCs from remediation processes to clean up contaminated soils and groundwater. Due to the focused regulatory attention on chlorinated organics, trichloroethylene (TCE) was chosen as a representative compound for this research.

#### 1.3 GAS PHASE ADSORPTION

Gas phase adsorption occurs on the surface and in the pores of the adsorbent. The adsorbate (the substance being adsorbed) is held by a combination of Van der Waals and electrostatic forces. The adsorption sites have a wide range of energy and adsorption occurs at the high energy sites first (Sontheimer et al 1988). In small pores the electrostatic fields from the adsorbent overlap and condensation of the gas into the pores is possible. This capillary condensation can dramatically increase the capacity of a given adsorbent. At higher contaminant concentrations (e.g. at higher partial pressures of contaminant, like water at high relative humidity) the adsorbent is more likely to form condensed films as the less energetic sites are filled. In addition, different molecules with different dipole moments and sizes will have different affinities for the various adsorbents.

The adsorbent capacities for the organic of interest are affected by several factors. At increased temperature the capacity is reduced. In fact, thermal processes are the primary way to regenerate adsorbents. Water will also compete for the adsorption sites and reduce the capacity of the adsorbent. A common way to reduce the relative humidity is to pre-heat the incoming air. This has been shown to be a cost-effective option on air stripping operations (Crittenden et al, 1988).

#### 1.4 CATALYSIS

Catalysis is a broad classification for reactions taking place where one of the participating species is not consumed or created, but rather can go on to continue the reaction. In combustion work it is primarily the thermal energy that activates the catalysts. The activated catalyst particles are usually dispersed on supports and the contaminated gas flows over them (similar to a catalyst bed in an automobile).

Other types of catalysis include photocatalysis where the light energy activates the catalyst or electrochemical catalysis (or electrocatalysis) where an applied potential aides in the reaction. Two areas this work focuses on is thermal catalysis and the enhancement (or depression) of the activity and/or selectivity of the catalyst by an external electric field.

In reactors where a catalyst surface is involved, like the car exhaust unit, a term called "heterogeneous" is used to describe the system. This is because the contaminant must diffuse from the bulk gas to the surface, adsorb onto the surface, react on the surface to create the desired end products, desorb the products from the surface, and, finally, diffuse the products away from the catalyst.

Using a carbonacaeous-based catalyst researchers (Petrosius and Drago, 1992, Petrosius et al, 1993) effectively destroyed methylene chloride and other chlorinated compounds at temperatures below 300 °C. They found that the activity of the catalyst was enhanced with the addition of transition metal (Mn, Co, Ti) oxides. This research extended their work to include the oxidation of trichloroethylene using the same base material and metal oxides.

### 1.5 ELECTRONIC FACTORS OF GAS PHASE ADSORPTION AND CATALYSIS

The phenomenon of adsorption and heterogenous catalysis rely on the molecule coming in contact with the surface. The nature of the electronic states within the surface and the molecules have direct bearing on the outcome of the collision. The electron cloud of the molecule can be distorted by the surface inducing a dipole within the molecule. As the molecule approaches the surface the potential energy of the molecule decreases due to the attraction of the mutually induced dipole moments. The distortion can be sufficient to allow a transfer of electrons, chemisorbing the molecule to the surface. This loss or gain of electrons by the adsorbate increases the reactivity of the adsorbed species. At surfaces with sufficient energy further reactions can occur with these radicals causing a change in the molecular form. The desired change is usually the loss of electrons from the parent molecule, oxidizing it to benign substances. Thus the electron, and its state, are key to these processes whether they are in the molecule or on the surface. With chemisorption of adsorbates over semiconductors, Vol'kenshtein (1963) described the process in terms of band bending. With a "clean" surface the bands of the semiconductor are the same in the bulk as on the surface. When a molecule chemisorbs to the surface, the energy band of the surface bends toward the band of the molecule. Each molecule adsorbed bends the bands an incremental amount until, at equilibrium, the bands are equal and adsorption ceases.

Holder (1970) evaluated the effect of an electrostatic field on the liquid-phase adsorption capacity of a bed of activated carbon. Using potentials from 0 to 25,000 V, he found a range of responses. The capacity of the activated carbon for urea was not altered, while the capacity for n-buteralydehyde increased by 113 percent at 16,000 V. Benzene and phenol capacities increased by 35 percent. Woodard et al. (1986) used flow through carbon fiber electrodes in aqueous streams to evaluate the electrosorption of various charged and neutral compounds. They found the adsorption process to be well described by a Freundlich isotherm and suggest using electrosorption for separating out small, functionalized organic molecules such as amino acids and proteins.

Gas phase electrosorption on semiconductors also occurs. Hoenig and Lane (1968) studied the oxygen uptake onto the n-type semiconductor ZnO. At +150 VDC there was little effect. At +300 V DC there was a drop in oxygen pressure, indicating adsorption onto the surface. Oxygen adsorption onto epitaxial  $\beta$ -HgS films under high fields (10<sup>5</sup> V/cm) was shown by Ivankiv and Muzychuk (1968). As the field increased to 4 x 10<sup>5</sup> V/cm, the conductivity increased monotonically until saturation of the surface occurred.

Adsorption itself can change the properties of the adsorbent. Researchers (McLintock and Orr, 1968; McIntosh et al. 1947) found that adsorption of gases onto carbon films and rods altered the bulk resistance of the materials. Others (Blackwood and Josowicz, 1991) showed the same effect for conductive polymers. The polymers contain conductive crystallites and non-conductive amorphous regions and the electrical conduction is by hopping from one crystallite to another, even up to 7 Å apart (Thakur and Elman, 1989). No information was available in the literature on the resistance changes upon adsorption for the carbonaceous adsorbents used in this research.

#### 1.6 JOULE HEATING

In electrically conductive materials the electron flow may be impeded by the molecules in the lattice. The energy loss due to this "friction" is converted to heat. The more electrons flowing (i.e. the larger the current) the more collisions and the more heat generated. This heat is generated within the matrix of the material and a bed heated in this manner is heated from the inside out. Irregular shapes and/or impurities in the adsorbent will give rise to preferential pathways in the bed, creating a potential for "hot" spots. Synthetic adsorbents of regular, uniform size, shape, and consistency should be ideal for minimizing these preferential pathways.

#### 1.7 OBJECTIVES

In this research three related areas were focused on to address the control of chlorinated organics from gaseous waste streams. The first was to look at the catalytic

oxidation properties of Ambersorb 572. It has surface areas similar to GAC, but is much stronger than GAC and can be regenerated in-situ. The material was used plain and impregnated with metal oxides of cobalt, titanium, and manganese.

The second area involved electric fields across two different adsorbents, Ambersorb 572 and an activated carbon cloth. The focus was on influencing the adsorption and catalysis reaction to increase capacity or lower the activation energy (operating temperature) of the catalysts, respectively. Dynamic (current flowing) conditions caused the adsorbent beds to heat up due to the Joule heating effect. The last area involved the resistance changes the activated carbon cloth undergoes upon adsorption of organic contaminants.

Specific objectives for this work include:

- A) Evaluating the Ambersorb 572 adsorbent, both plain and impregnated, as catalysts in the oxidation of TCE at temperatures below 250 °C.
- B) The impact of electric fields on adsorption on activated carbon cloth, and the impact of electric fields on the catalytic reactions over Ambersorb 572 (plain and impregnated).
- C) Determine the resistance change upon adsorption on the carbon cloth for seven compounds. Evaluate the impact of concentration on the resistance change for trichloroethylene on the carbon cloth.
  - D) Field demonstrate a lab-scale system at a contaminated Air Force site.

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#### Chapter 2

Catalytic Oxidation of Trichloroethylene over Carbonaceous Adsorbents

#### 2.1 INTRODUCTION

Industrial and remediation processes generate large quantities of air contaminated with organics. An estimated 1.22 billion pounds of organics are released to the air each year from U.S. facilities (USEPA, 1993). Clean Air regulations promulgated by the U.S. Environmental Protection Agency (EPA) place limits on the types and amounts of organics that can be released to the environment. Chlorinated organics, specifically trichloroethylene (TCE), were targeted for destruction in this study because of their suspected toxicity and because of the large quantity of TCE being released to the atmosphere (in the top ten, based on quantity released, on the USEPA, 1993 list).

Technologies currently used to limit the release of the organics to the air include gas phase adsorption onto granular activated carbon (GAC) and combustion, either in a direct flame incinerator at high temperatures (>2000 °C) or over a catalyst at moderate temperatures (200-500 °C). An example of a catalytic unit is one evaluated by Hylton (1992). It utilizes a chromia oxide catalyst in a fluidized bed and operates at an Empty Bed Contact Time (EBCT) of 0.5 s at 370 °C to achieve a 97 percent conversion of trichloroethylene (TCE). Higher temperatures and/or longer time in the reactor (residence time) allows for complete conversion or mineralization of the chlorinated organics to water and mineral acids such as HCl. Partial conversion of the molecules can result in products that are more toxic than the parent compound (e.g. phosgene, COCl<sub>2</sub>). By-products of combustion are unwanted but may be tolerable if they are benign.

Destruction of gaseous by-products from automobile exhaust fueled much of the early catalysis research, leading to catalyst beds with mixtures of precious metals finely dispersed on high surface area ceramic supports. High surface area polymer catalysts are

important in industry, especially the fuel additives industry (Al-Jarallah et al, 1988). However, the focus on using them for catalytic oxidation reactions to mineralize organic pollutants has only recently been reported (Petrosius and Drago, 1992, Petrosius et al, 1993).

Petrosius and Drago (1992) evaluated a number of metal oxides on Ambersorb® adsorbent supports as well as Ambersorb 563 & 572 alone for oxidation of methylene chloride (250 °C, EBCT=240 s for all experiments). While the supports alone are quite active (68 and 78 percent conversion), a chromium oxide catalyst impregnated onto the surface increases the conversion to 99.9 percent. Titanium dioxide (TiO<sub>2</sub>) on Ambersorb 572 showed a 93 percent conversion and a cobalt oxide catalyst also on Ambersorb 572 showed a 75 percent conversion.

In subsequent work the same group (Petrosius et al, 1993) reported the decomposition of several halogenated hydrocarbons over the same types of supports/catalysts at the same conditions (250 °C, EBCT=240 s). They found a 16 percent conversion of tetrachloroethylene (PCE) in humidified air over a chromia oxide impregnated Ambersorb 572 catalyst/support. On plain Ambersorb 563, carbon tetrachloride (CCl<sub>4</sub>) conversion in dry air dropped from 73 to 5 percent in 120 hours. By adding water the conversion rose to 47 percent at the 170 hour mark. In other experiments they also noted a 67 percent decrease in the conversion percentage of methylene chloride when the flow was increased by a factor of 5 (EBCT from 240 s to 49 s) with dry feed at 250 °C.

Since the catalytic oxidation process involves electron exchange, it was thought that an applied electric field would alter the process, possibly reducing the activation energy of the reaction. This would in turn reduce the thermal energy requirements for a given conversion. Dynamic fields (current flowing) were passed through beds of reactive Ambersorb 572 to test this hypothesis.

In this research an evaluation was made on the impact of operating the catalysts at low temperatures and short residence times on the catalytic oxidation of TCE from gas streams using transition metal impregnated and non-impregnated Ambersorb 572 adsorbent/catalysts. In addition, the impact of an applied electric field was also evaluated. The impact of the flow rate, catalyst type, and current type (AC or DC) was also evaluated.

#### 2.2 MATERIALS AND METHODS

All chemicals used in the experiments and analyses were reagent grade or better. Reagent grade dichloromethane (99.9+%), titanium (IV) chloride (99.995+%), sodium persulfate, methanol, sodium hydroxide, sulfuric acid, cobalt and manganese nitrates, and potassium permanganate, were purchased from Aldrich Chemical Company (Milwaukee, WI). Water for the bubblers came from a Milli-Q® station (Millipore, Molsheim, France) in the lab. TCE in dry nitrogen (500 parts per million by volume [ppmv]) was purchased from Scott Specialty Gas Company (Riverside, CA). TCE in dry air (15 & 1500 ppmv) were purchased from Matheson Gas Products (Jolliet, IL). The Ambersorb 572 adsorbent (40x60 mesh, Lot No:923318) was purchased from the Rohm and Haas Company (Philadelphia, PA). All carrier gases including air, argon, helium, nitrogen were ultra-pure and purchased from Interstate Gas Company (Marinette, MI).

On-line gas-phase samples were taken every two hours by an automated sampler and analyzed according to the Environmental Protection Agency's Reference Method 23 (Scott Environmental, 1983) using a Hewlett Packard Model 5890 Series II gas chromatograph equipped with a flame ionization detector (FID) and an electron capture detector (ECD). Compound separation was performed using a stainless steel VOCOL<sup>TM</sup> (Supelco, PA) column 60 m long by 0.75 mm diameter with a 1.5  $\mu$ m film. The gas samples passed through heated stainless steel lines into a six-port valve (Model RD6P, Valco) that was heated to 160 °C (Valco Instrumentation Temperature Controller). A 0.5 cm<sup>3</sup> sample loop attached to the six-port valve provided a gas sample size that yielded a TCE detection limit of about 1 ppm using the FID and about 10 ppb with the ECD. All

reaction efficiencies reported in this study are steady-state conversions, unless otherwise noted.

Two bubblers were used to trap chlorine and sulfur compounds after the 6-port valve. The first in series contained Milli-Q<sup>®</sup> water to trap chloride and sulfate ions and the second trap had sodium persulfate to trap chlorine gas. Chloride and sulfate ions were analyzed using an isocratic elution of anions with a 21mM NaOH solution (flowrate 1ml/min), on a Dionex DX-500 Ion Chromatograph (Dionex Corp., Sunnyvale, CA) equipped with an Ionpac AG11 guard column (4 mm), an Ionpac AS11 analytical column (4 mm), and a Dionex CD20 conductivity detector.

Periodic effluent samples from the gas-phase catalytic reactor were taken to identify by-products, especially chlorinated species not trapped by the water/sodium persulfate bubblers. These were bubbled through dichloromethane (DCM) and the DCM was then analyzed with a Hewlett Packard Model 5970B mass selective detector with electron impact ionization controlled by a Hewlett Packard 5990C Chemstation data system (Hewlett Packard, Minneapolis, MN). A DB1701 (14% cyanopropylphenylmethylpolysiloxane) capillary column (30 m length, 0.25 mm ID, narrow bore, 0.25 micron film thickness) was used to separate the compounds (J&W Scientific, Folsom, CA). One milliliter per minute of helium was used as the carrier gas. From an initial temperature of 35 °C, the oven temperature was ramped to 230 °C at a rate of 5 °C/min and kept there for a period of six minutes. The injection port and detector temperatures were maintained at 250 °C. The total ion chromatograph plot was generated by scanning masses between 30-800 amu. The fluid from the methanol (solvent) extraction of the exposed catalyst (MTU-CAT) was also analyzed on the HP 5970B under the same program conditions.

#### 2.2.1 Catalyst Impregnation Procedures

The following procedures were used to impregnate oxides of cobalt, manganese and titanium onto plain Ambersorb 572 (AM 572-P). Impregnation of these

metal oxides onto the AM 572-P was carried out using the incipient wetness technique outlined by Petrosius et al. (1993) except as noted below. Solutions of cobalt and manganese nitrate salts or TiCl<sub>4</sub> were added separately to 100 gram lots of the adsorbent. The catalysts were loaded to 14 percent by weight metal/dry gram of adsorbent. The solutions were added from a buret drop-wise until a volume equal to the pore volume of the resin was added. Water and dichloromethane were used to make up the remaining volume requirements for the cobalt/manganese nitrates and TiCl<sub>4</sub> solutions, respectively. The impregnated resin was wet in appearance but there was no separate liquid phase. The titanium tetrachloride impregnated adsorbent was allowed to react with the room humidity to form titanium dioxide. An X-ray diffraction of TiO<sub>2</sub> residue mixed the same way, but allowed to evaporate in a glass vial, indicated over 95 percent of the material was in the anatase crystal form. Therefore, we assume that the TiO<sub>2</sub> on the impregnated adsorbent is anatase which is thought to be more catalytically active. The other materials were heated overnight at 180 °C in an oven (Isotemp Oven Model 655G, Fisher Scientific) under a vacuum supplied by a pump external to the oven. The Co and Mn impregnated adsorbents were then heated in separate batches under nitrogen at 360 °C for 12 hours to remove the nitrates. The catalysts were transferred to 125 ml Erlenmeyer flasks and capped until used.

#### 2.2.2 Experimental Reactor Design

The experimental set-up is displayed in Figure 2-1. TCE in air was metered from pressurized tanks through the reactor followed by a chloride ion trap and exhausted to a fume hood. An optional water bubbler at room temperature was placed before the reactor to provide a humidified air stream (Relative Humidity, R.H., was about 45 percent). Valves were placed before and after the reactor to monitor the TCE concentration. The reactor was wrapped with heat tape to control the temperature of the reactor. A small thermocouple was placed inside the reactor, near the middle of the bed to monitor the temperature. A second thermocouple was placed outside the bed, under the heat tape wrapping to monitor the exterior temperature. Two bubblers were placed at the end of the effluent line, one containing Milli-Q® water for trapping chloride and sulfate

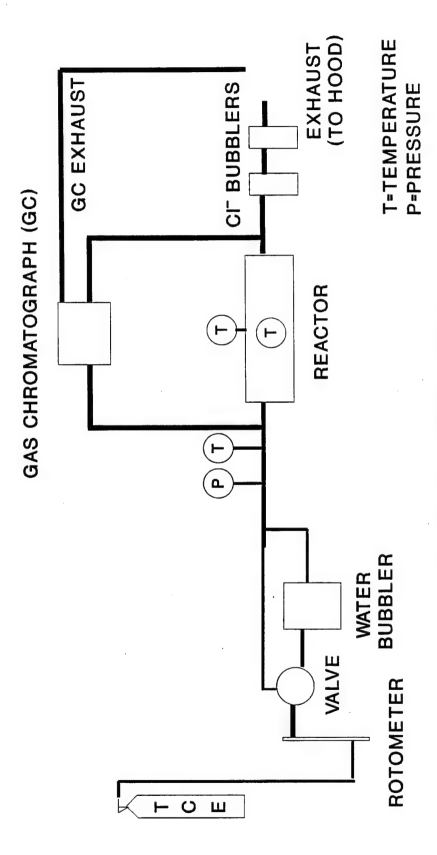


Figure 2-1. System Setup

ions, and the other containing saturated sodium persulfate solution for trapping the chlorine gas. The packed-bed reactor consisted of either a 1.5 inch I.D. by 6 inches long carbon steel column or a glass tube (1/4 inch OD; 1/8 inch ID). Due to the adsorption potential of the catalyst support, some experiments were conducted with materials that were loaded to capacity at room temperature. These are referred to as "pre-loaded" adsorbent/catalysts in this paper.

The oxidation reactor, shown in expanded form in Figure 2-2, was used for all experiments - thermal only and those with an applied electric field. The graphite rod (The Carbide/Graphite Group, Inc., St. Marys, PA; no part or lot number) in the center of the reactor was 1/8 inch in diameter and extended to within 1/2 inch of the bottom. A section of conductive, activated carbon cloth ("Kuractive," FT 200-15, Lot # FK 622096-1B) was used in the bottom of the reactor to keep the adsorbent/catalyst from falling out of the reactor. Electrical isolation was accomplished with Teflon® tape and glass tubing connectors.

All the gas flow rates reported herein are based on 1 atm and 22 °C unless otherwise stated. The empty bed contact times (EBCT = volume of catalyst bed/volumetric flow rate) or the space velocities (EBCT<sup>-1</sup>) are based on the flow rate at 22 °C. The pressure drop across the bed at the different flowrates used in this study changed by less than 0.5 psi and did not significantly alter the vapor phase concentration.

The term conversion is determined by the amount of TCE in the exit stream as compared to the amount in the feed stream as indicated by the GC. A 100 percent conversion is defined as the complete loss of the TCE in the effluent. Mineralization is defined as the amount of HCl or Cl<sub>2</sub> found in the traps as compared to that which is expected from stoichiometry. A 100 percent mineralization level indicates that all of the available incoming chlorine is accounted for in the traps. Sometimes TCE that is preadsorbed on the adsorbent/support will come off and be mineralized in addition to that being fed to the reactor. Under these conditions it is possible to have a mineralization percentage greater than 100 percent.

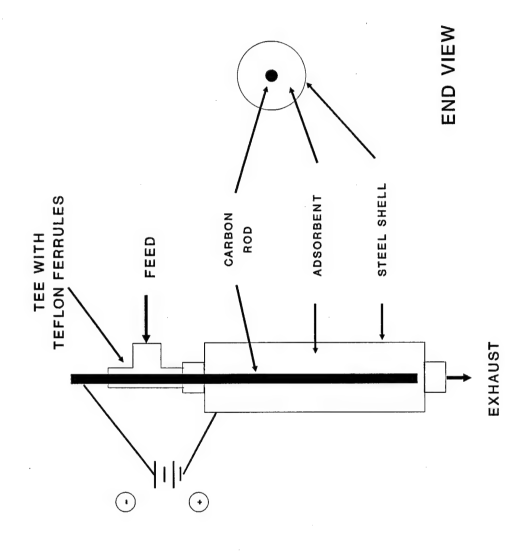


Figure 2-2. Annular Reactor Design

Catalyst support characterizations were evaluated by observing the changes in the surface area as determined by BET analysis and also by direct elemental analysis of the material before and after exposure to the TCE stream to be treated. These support analyses were performed by Porous Materials, Inc. (Ithaca NY) and Huffman Laboratories, Inc. (Golden CO), respectively.

#### 2.2.3 Ambersorb 572 Characteristics

The plain Ambersorb adsorbent (AM-572-P) has the characteristics shown in Table 2-1. The material is a highly sulfonated styrene divinylbenzene macroreticular ion exchange resin that is pyrolyzed in a patented process (Rohm and Haas, 1992). An elemental analysis revealed that the material is 1.45 weight percent (± 0.1 percent) sulfur. If each sulfur atom forms a -SO<sub>3</sub>H group, and if there is a three-dimensional node of seven -SO<sub>3</sub>H groups forming one catalyst site (Wesley and Gates, 1974), then there are approximately 4 x 10<sup>19</sup> sites per gram of material. Turnover numbers (moles reacted per minute/mole of catalyst sites) ranged from 0.2 to 0.002 in this study.

TABLE 2-1. PHYSICAL PROPERTIES OF AMBERSORB 572<sup>†</sup>

Surface Area m²/g	Microporosity (<20 A) ml/g	Mesoporosity (20-500 A) ml/g	Macroporosity (>500 A) ml/g	Particle Size	Bulk Density g/cc
1100	0.41	0.19	0.24	20 to 50	0.49

<sup>† -</sup> As reported by Rohm and Haas (1992).

#### 2.2.4 Formation and Characterization of MTU-CAT

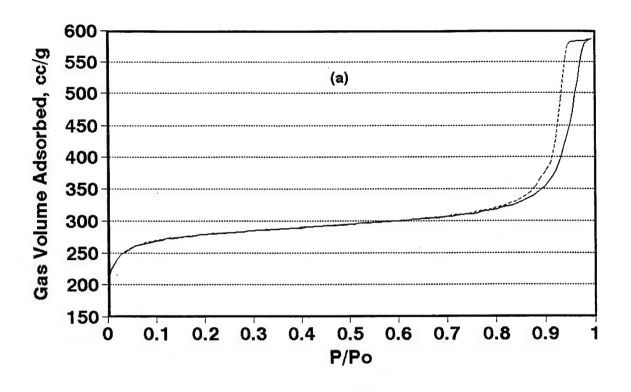
While using an experimental heating process the Ambersorb 572 base material was converted into MTU-CAT. The heating process utilizes electrical energy to resistively (Joule) heat the adsorbent. As the current flows through the bead particles the

electrons impact the lattice of the adsorbent heating it. The current flows around the pores and the actual electron path inside a bead may be quite tortuous. If large currents are used (e.g. using 1.3 Amps/cm² to make MTU-CAT) then areas inside the particles with small cross sections may carry a large current load. This current load may be sufficient to burn away portions of the material, even though the bulk bed temperature remains below the thermal limits of the material.

Evidence for this morphological change in the MTU-CAT material is indicated by the adsorption/desorption BET analysis data shown in Figure 2-3. The hysterisis loop in the plain Ambersorb 572 is a Brunauer, Deming, Deming, and Teller (BDDT) Type IVa isotherm which is indicative that the material posses straight walled pores (e.g. vertical wells) as the pore shape (Gregg and Sing, 1982). The MTU-CAT shows that these vertical wells still exist, but the separation of the adsorption and desorption lines suggest that slits are formed in the material, similar to a BDDT Type IVb isotherm. One possible way for the slits to form is the electrolytic melting of the material between two adjacent wells. Another possibility is the physical separation of the graphitic planes within the polymer matrix. Additional catalyst sites, differing in performance from the plain material, may have been created in the slits.

#### 2.3 RESULTS AND DISCUSSION

The polymer matrix supports -SO<sub>3</sub>H groups on the surface that form acid catalyst sites. These sites can be ion exchanged and the rate of the acid-catalyzed reaction is dependent on the concentration of the -SO<sub>3</sub>H sites. When the -SO<sub>3</sub>H population is dense then they form clusters. These clusters then behave in a manner similar to catalysis in concentrated acid solution. The polymer would then be able to solvate and stabilize reactive intermediates such as carbenium ions, with the following suggested structure (Wesley and Gates, 1974):



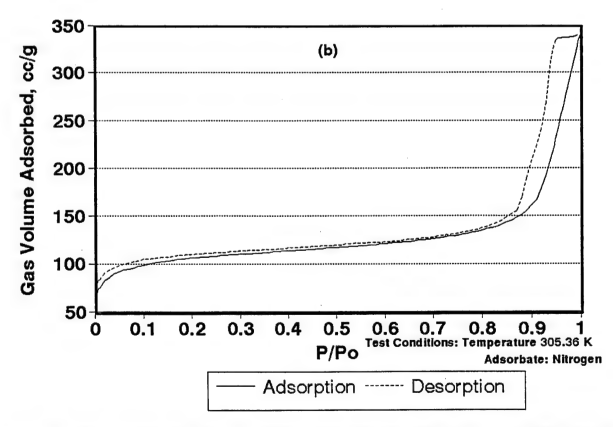
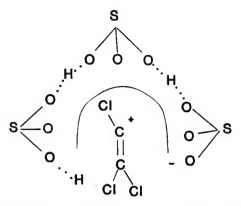


Figure 2-3. Gas Volume vs P/Po: (a) Virgin AM572-P; (b) MTU-CAT



The rate of reaction will be dependent on the number of catalyst sites. Two main factors that would affect the concentration of reactive species at these sites are temperature and the localized TCE concentration. If the temperature is too low then very few reactive species will be formed and coking of the catalyst may occur. If the concentration of TCE is too low in the pores with the capabilities of supporting the reactive species then only a few reactive sites would be formed. If the temperature is too great then the reactive species concentration on the surface will go down. Thus it is possible that there is an optimum temperature to achieve the highest capacity for the reactive species on the surface. This optimum temperature would be dependent on the feed concentration as well as the organic being destroyed because the surface concentration is dependent on the type of organic and its concentration.

The structure of the polymeric adsorbent will dictate how many catalyst sites will be formed and the reactions that occur on them. The -SO<sub>3</sub>H functional groups must be close enough to interact and form the stabilized center for the reactive intermediate. The rigidity of the polymer dictates how these functional groups are held. If the polymer is too flexible it does not efficiently create the stabilized centers, and fewer catalyst sites are formed. Once formed they are reactive and polar molecules (e.g. water) have a high affinity for the -SO<sub>3</sub>H groups and exclude reactants and suppress the catalytic activity (Gates, 1992).

Due to the adsorptive nature of the catalyst support, the importance of the adsorption of reactants and products on the surface cannot be over emphasized. Ambersorb 572 was found to have an adsorption capacity of 0.17 ml TCE/g adsorbent at 196 °C when fed a

dry feed stream of TCE (~80,000 ppmv in nitrogen). For this reason chloride analysis of the product stream is the only way to confirm mineralization. The amount of chloride also gives an indication of how the catalyst is performing over time. There is a general decreasing trend in the chlorine content of the effluent, possibly the result of catalyst poisoning with age.

The oxidation of TCE can occur with or without water according to the following reactions:

$$2 C_2HCl_3 + 4\frac{1}{2} O_2 --> 4 CO_2 + 6 Cl_2 + H_2O$$
  $\Delta G_{200^{\circ}C} = -223 \text{ kcal/gmole TCE}$ 

$$C_2HCl_3 + H_2O + 1\frac{1}{2}O_2 --> 2 CO_2 + 3 HCl$$
  $\Delta G_{200^{\circ}C} = -216 \text{ kcal/gmole TCE}$ 

The conversion of the plain support alone, AM 572-P, was substantial (>99.9 percent at 180 °C). The conversion of the impregnated materials ranged from none to complete conversion. The designations of the catalysts are AM-572-X, where X is the metal oxide code: Mn for manganese; Co for cobalt; and Ti for titanium. Experiments conducted on a thermally modified non-impregnated Ambersorb 572, referred to as MTU-CAT, showed even higher conversion and mineralization levels than the plain support. The following sections outline the experiments of these catalysts. Catalyst aliquots were used in either the "pre-loaded" form or the "as-received" form in the following experiments.

#### 2.3.1 AM-572-P

Experiments using the pre-loaded form (see Table 2-2), were initially performed at 130 °C with no conversion detected. Increasing the EBCT from 1.7 s to 2.7 s by decreasing the gas flow rate did not change the conversion significantly, however chloroform, carbon tetrachloride, and PCE, were detected in the effluent. Increasing the temperature to 180 °C at the same EBCT caused the effluent to drop to below detection limits (>99.9 percent conversion), yet the chloride balance only showed 20 percent Cl

recovery. After 380 hours (approximately 1 gram TCE fed/g catalyst) the catalyst was operating at only 50 percent conversion.

Performance of the as-received (not pre-loaded) catalyst was drastically different. In these experiments the catalyst was maintained at 180 °C at all times. Flow rates varied from 1.1 s to 16.7 s EBCT. Water addition (50 percent RH) to the feed stream had no effect on the catalyst performance. This is a surprising finding since the water reacts so strongly with the polarized -SO<sub>3</sub>H catalysis groups. Therefore, either the groups are occupied with something else or they are not responsible for the oxidation occurring on this catalyst.

TABLE 2-2. OPERATING CHARACTERISTICS OF AM-572-P EXPERIMENTS (1500 ppmv TCE feed).

Temperature (°C)	EBCT (s)	Conversion (%)	Mineralization (%)	Water (Y/N)
		PRELOADED		
130	2.7	0	NT	N
180	2.7	>99.9	20	N
		AS RECEIVED		
180	16.7	>99.9	70	N
180	16.7	>99.9	70	Y
180	1.1	>99.9	NT	N
180	0.5	97	NT	Y
180 <sup>†</sup>	0.15	20	34	Y
180 <sup>†</sup>	0.15	10	14 <sup>a</sup>	Y

NT - Not Taken

Tabled data is a subset of 192 samples.

Since adsorption is possible, the catalyst was operated until approximately 0.8 g of TCE/g catalyst passed through the system. Chloride samples confirmed the

<sup>&</sup>lt;sup>a</sup> After 3 days.

<sup>&</sup>lt;sup>†</sup> Concentration switched to 15 ppmv.

destruction of TCE as mineralization efficiencies remained high (approximately 86 percent at the end of the experimental sequence). A minimum turnover for the plain, as-received catalyst is approximately 8 x 10<sup>5</sup>/ mole of catalyst sites. It is possible that as the preloaded catalyst heated up, the catalyst sites were exposed to TCE at less than optimum temperatures and fouled by coking of the sites.

### 2.3.2 MTU-CAT

Table 2-3 summarizes the catalytic oxidation experiments of TCE over the MTU-CAT for a wide range of operating conditions. No "pre-loaded" experiments were performed with MTU-CAT. Exposing MTU-CAT to a dry, 1500 ppmv TCE feed, at a catalyst temperature of 200 °C and an empty bed contact time (EBCT) of 7.7 sec (17.5 ml/min-g catalyst) revealed a very active catalyst. The effluent, as determined by the Flame Ionization Detector (FID) on the GC, remained non-detect for TCE (99.9 percent conversion) and a small amount (approximately 1 ppm) of carbon tetrachloride was detected in the effluent. Approximately 3 g of TCE was fed to the reactor for each gram of catalyst during these initial experiments shown in Table 2-3. The same adsorbent/ catalyst charge was used throughout the series of conditions listed in Table 2-3. Some of these early MTU-CAT experiments only used a water bubbler to trap HCl/Cl<sub>2</sub>. All remaining experiments reported in this study used an additional sodium persulfate trap for the Cl<sub>2</sub> specifically and increased the chlorine recovery efficiency in the exhaust. Table 2-3 annotates which type of trapping system was used in the initial MTU-CAT experiments. The shift in the effluent from all HCl to greater amounts of Cl<sub>2</sub> suggests the initial reactions reacted with residual water on the adsorbent (desorption studies at 110 °C show approximately 30 mg water/g adsorbent in the "as received" form). As the available water decreased, the reaction shifted to a true dry reaction sequence, but this formed a small amount of water for the "humid feed" reaction to continue.

Additional tests were performed on subsets of the MTU-CAT used in the above experiments. In one experiment (140 °C, 2 s EBCT), a dry TCE feed was used and the effluent dropped to below detection limits on the very sensitive Electronic Capture

Detector (ECD) (>99.999 percent conversion). Analysis of a DCM sample taken while using the dry feed indicated that phosgene, and other by-products (e.g. PCE, hexachloroethane, and possibly some diethoxy ethanes) were being formed in trace amounts.

Water was then added to the dry feed to produce a TCE/air mix with a relative humidity of 45 percent at 22 °C. Within minutes this humidified feed rapidly deactivated the catalyst. Upon removing the water, the catalyst performance slowly recovered. This is in direct contrast to the AM 572-P which showed no effect of added water. This

TABLE 2-3. OPERATING CHARACTERISTICS OF THE INITIAL DRY MTU-CAT EXPERIMENTS (1500 ppmy TCE feed - no water added).

Temperature	Empty Bed	Conversion	Mineralization	Time
(°C)	Contact Time	(%)	$(\%)^{\ddagger}$	(hr)
	(s)			
200	7.7	>99.9	104	70
200	7.7	>99.9	80	90
180	24.5	>99.9	127 <sup>†</sup>	
180	9.8	>99.9	NT	
180	4.8	>99.9	NT	
180	3.0	>99.9	62	
150	2.7	>99.9	90	
115	2.9	>99.9	52	190
115	2.9	>99.9	56/(73)	214
110	11.7	>99.9	37/(59)	260

NT - Not Taken

Tabled data are a subset of 132 samples.

<sup>&</sup>lt;sup>†</sup> Additional processes evaluated with these experimental runs caused an excess amount of TCE to be desorbed from the adsorbent adding to the chlorine balance.

<sup>&</sup>lt;sup>‡</sup> Dual trap values are shown in parentheses

finding indicates the catalyst sites or the catalytically active species are different from those in the plain material. A minimum turnover number for the dry feed reactions is approximately 8 x 10<sup>5</sup>/ mole of catalyst sites, coincidently the same as the AM572-P.

In separate experiments the addition of natural gas (CH<sub>4</sub>, 180 °C, concentration approximately 0.15 percent) to the dry TCE/air mix caused a dramatic drop in the TCE conversion. The catalyst performance returned after removal of the natural gas. The formation of water by the oxidation of methane is possible, accounting for the loss in conversion.

BET area analyses, elemental analyses, and a liquid extraction technique were used to elucidate the mechanisms of catalysis and to determine the structural differences of this enhanced catalyst, MTU-CAT. The surface area of MTU-CAT has changed from the parent Ambersorb 572 adsorbent as shown in Table 2-4. There is a shift in the distribution of the pore sizes indicating that the material has indeed changed. However, it is unclear if the change is due to pore blocking/filling from by-products or if the change occurred during the electrolytic activation of the MTU-CAT. Figure 2-4 graphically shows the results of the nitrogen isotherm data.

TABLE 2-4. SURFACE AREA ANALYSIS RESULTS FOR VIRGIN AMBERSORB 572 AND MTU-CAT AFTER THE 11 DAY INITIAL EXPERIMENTS.

SAMPLE ID	Total Surface Area (m²/g)	Average Pore Diameter (Å)	Total Pore Volume (ml/g)	Median Pore Diameter (Å) <sup>†</sup>	Median Pore Diameter (Å) <sup>‡</sup>
572-P	831	44	0.91	132	56
MTU- CAT	324	65	0.53	296	64

Analysis conditions: Instrument Temperature, 305.36 °K; Room Temperature, 296.7 °K; Adsorbate, Nitrogen.

<sup>†</sup> Based on Pore Volume; † Based on Surface Area

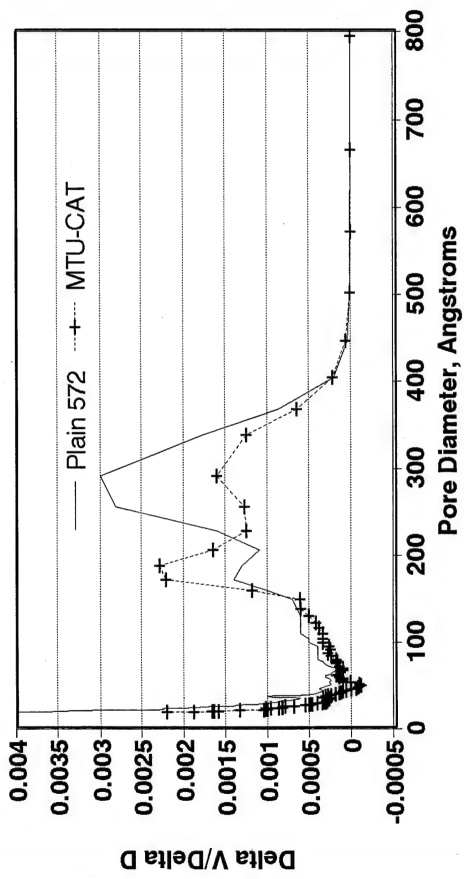


Figure 2-4. BET Adsorption/Desorption Isotherm Data; Delta V/Delta D vs Pore Diameter

The results of elemental analyses on the parent compound (AM-572-P), the enhanced catalyst (MTU-CAT), and a methanol extracted sample of the enhanced catalyst are shown in Figure 2-5. They indicate a dramatic chlorine increase in or on the MTU-CAT. At least some of the chlorine is leachable, as the methanol extracted sample has lost some of the chlorine residual. A GC/MS examination was made of the extracted liquid to identify these compounds (listed in Table 2-5) leached from the catalyst.

TABLE 2-5. GC/MS ANALYSIS OF METHANOL EXTRACT OF MTU-CAT.

Compound	Relative Peak Area
	(Percentage)
Hexachloroethane	12.4
Pentachloroethane	9.25
Tetrachloroethylene (PCE)	2.54
Trichloroacetic Acid, Methyl Ester <sup>†</sup>	1.47
1,1,2,3,4,4, hexachloro, 1,3 Butadiene	0.39
Dichloroacetic Acid, Methyl Ester <sup>†</sup>	0.29

<sup>&</sup>lt;sup>†</sup> The methylated ester of the parent acid was probably created when the methanol reacted with the acid over the -SO<sub>3</sub>H groups (Morrison and Boyd, 1980).

The existence of the dichloroacetic and trichloroacetic acids on the catalyst surface, and the phosgene noted earlier, suggest a radical mechanism similar to the photocatalysis of TCE in the gas phase over titanium dioxide. Nimlos et al (1993) identified these compounds as by-products when the hydroxyl radical (•OH) attacks the TCE. They report that the reactive chlorine atom, •Cl, is formed, then this in-turn attacks the organics to make the dichloroacetic acid. In addition, they suggest that the same reaction mechanisims would occur if PCE was the target organic instead of TCE. Under these conditions trichloroacetic acid would be formed.

It is also possible that the radical in this catalyst could be  ${}^{\bullet}C_2Cl_3$  based on the recovery of 1,1,2,3,4,4 hexachloro butadiene (radical:radical recombination) from the surface. Other species could also exist, such as the carbenium ion  $C_2H_2Cl_3+$ , that could be

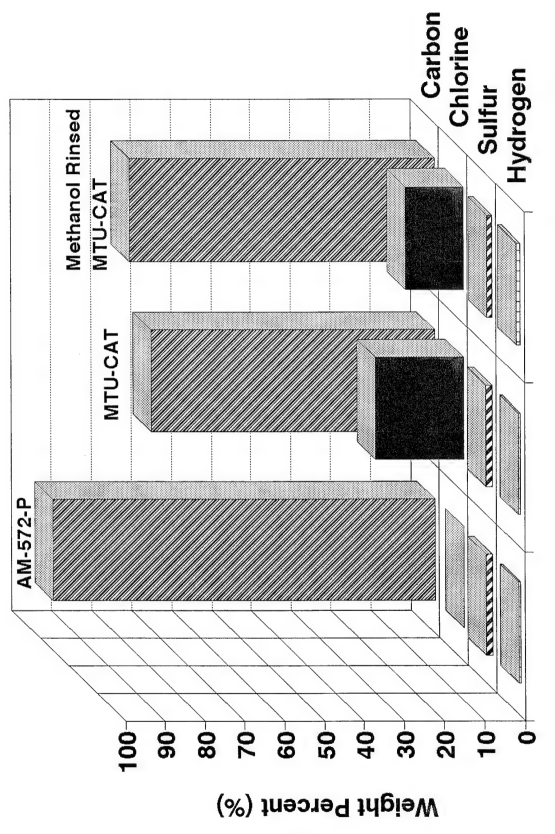


Figure 2-5. Elemental Analysis of the Materials used in the Oxidation of TCE

stabilized by the sulfonate groups on the polymeric support. No matter how these surface adsorbed by-products are created, they do increase the chlorine loading of the catalyst. These and potentially other polymerized, chlorinated by-products, are on the catalyst surface.

It was expected that the hydrogen levels would be depleted in the catalyst due to the hydrogen interactions noted earlier, however there is no statistical difference between the amount of hydrogen in the samples. This suggests that the hydrogen is protected within the matrix of the polymer and not exposed to the reaction sequence. Unfortunately the same cannot be said of the carbon matrix itself as there is a loss of carbon in the MTU-CAT material. It is unclear if this is due to the structural damage noted above (which may account for some of the loss in surface area). Whatever the cause, the enhanced catalyst has lost the volume equal to that contained in the micropores of Ambersorb 572 (0.4 ml, [Rohm and Haas, 1992]). If the micropores are blocked off, this suggests that the -SO<sub>3</sub>H groups in the mesopores are in sufficient proximity and numbers to support the reactive intermediate species (e.g., carbenium ion/radical) to oxidize TCE. However, Figure 2-4 indicates that only about two thirds of the micropores are lost.

#### 2.3.3 Metal-Impregnated Ambersorb 572

The experimental results with AM-572-Mn and AM-572-Co, shown in Table 2-6, were conducted with a dry feed. As shown in Figure 2-6, the catalysts appear to have induction times before they become active. This possibly occurs in two steps. First, is the loading of the high-energy sites in the pores (adsorption). Second, building up concentrations of reactive intermediates supported in the -SO<sub>3</sub>H groups in the pores. Once a critical loading of reactive intermediates occurs, then the reaction consumes the excess TCE shown exiting the reactor in Figure 2-6, in addition to removing the influent TCE. Water shows a fouling trend in both of these catalysts as well indicated by the increase in TCE concentration shown in Figure 2-6. No chloride samples were taken to determine mineralization efficiencies, nor were the experiments continued to evaluate the

reversibility of the water poisoning. The AM 572-Mn experiments (210 and 180 °C at 3.8 and 7.6 s EBCT) which started with humid feeds showed little conversion (0-12 percent) as shown in Table 2-6, again showing the deactivation effect water has on the catalysts.

TABLE 2-6. OPERATING CHARACTERISTICS OF COBALT (-Co) AND MANGANESE (-Mn) IMPREGNATED AMBERSORB 572

(1500 ppmv TCE feed).						
CATALYST	Temperature (°C)	EBCT (s)	Conversion (%)	Mineralization (%)	Water (Y/N)	
-Mn	175	16.9	>99.9ª	NT	N	
	175	16.9	80-88	NT	Y	
	210	3.8	0	NT	Y	
	210	7.6	12	NT	Y	
	180	7.6	5	NT	Y	
-Co	140	6.7	0	NT	N	
	140	10.3	0	NT	N	
	140	49	0	NT	N	
	190	24	80	17	N	
	190	24	>99.9	NT	N	
	190	9.6	>99.9	47	N	
	190	2.8	>99.9	NT	N	
	130	32	>99.9	27	N	
	130	9.6	>99.9	20	Ń	
	130	9.6	99.9	NT	N	
	130	3.0	40	NT	N	

<sup>&</sup>lt;sup>a</sup> After induction period - see text.

Tabled data are a subset of 102 samples (Mn) and 133 samples (Co).

Heat treatment of the cobalt impregnated catalyst was done to ensure the oxides were sufficiently formed prior to exposure to the TCE. The heating was done by taking an aliquot of the initial 100 g batch of AM572-Co and placing it in a tube furnace

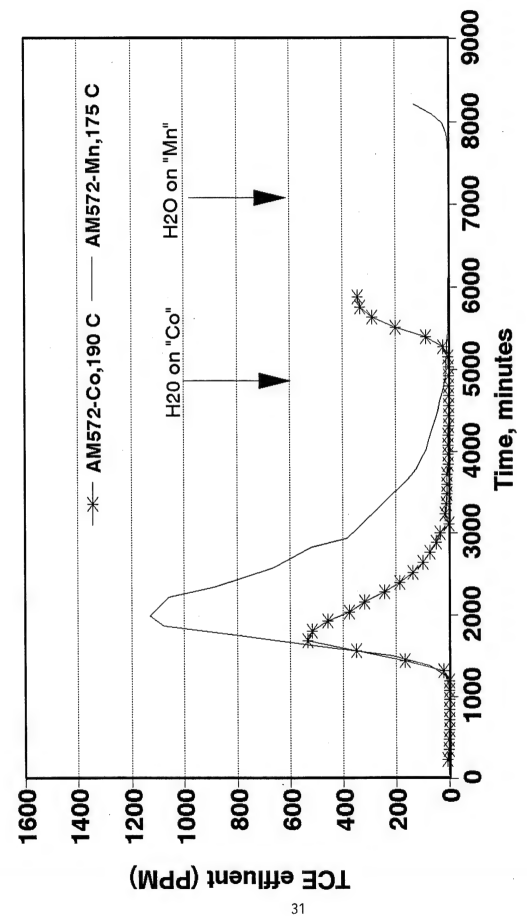


Figure 2-6. Organic Concentration Vs Time; EBCT=17 s, 1 atm, 1500 ppm TCE feed

and heating to 250 °C in air over night. Some of this material, AM572-Co(H), was put into the steel reactor (25 s EBCT; 280 °C). Only a 67% conversion was obtained, with chloroform and PCE as the major by-products. The catalyst suffered a 12 percent weight loss during the experiment.

In an attempt to specifically target the oxides onto the sulfonic functional groups, cobalt was ion exchanged with the hydrogen in the -SO<sub>3</sub>H groups. This new batch of catalyst was made by passing cobalt nitrate (1 M) through a bed of plain Ambersorb 572. As expected the rigid, pyrolyzed adsorbent has a low exchange capacity (less than 1/6th that of IRN 77) because of the inability of the adsorbent to swell to accommodate the larger cobalt cation. This catalyst, AM 572-Co(e), at 180 °C (1 s EBCT) also had an induction period, after which complete conversion (>99.9 percent) is achieved, but the mineralization is only 7 percent. Shortly afterwards the catalyst completely deactivates. Heat treating an aliquot of the AM-572-Co(e) using the same procedures as the AM572-Co above to make AM 572-Co(e)/H resulted in a non-active catalyst. The experiments showed a typical adsorption curve (short loading with breakthrough) and no conversion.

The results of experiments with a pre-loaded AM-572-Ti are shown in Table 2-7. Early high mineralization efficiencies may be the result of excess TiCl<sub>4</sub> conversion and not destruction of TCE. Adding Milli-Q water to the AM-572-Ti in a glass vial causes it to get very hot, confirming the existence of residual TiCl<sub>4</sub> in the pores. To remove this effect the catalyst, referred to as AM-572-Ti(W), was washed, dried, and preloaded before use. As the catalyst heated up to 200 °C, approximately 1/3 of the preloaded TCE exited the reactor along with other by-products, including PCE. Other by-products detected (typically found in this work) during the remainder of the experiments with this catalyst include carbon tetrachloride, chloroform, hexachloroethane and possibly pentachloroethane. The product spectrum shifted over time with the hexachloroethane concentration decreasing while the TCE concentration in the effluent was increasing. These excessive by-products do not favor the use of AM-572-Ti.

The product spectrum between the metal-impregnated and the plain catalysts are similar. In addition, there does not seem to be any enhancement of the catalyst performance with the addition of the metal oxides. It is possible that these temperatures are too low to activate the reactions over these metal oxides.

TABLE 2-7. OPERATING CHARACTERISTICS OF TITANIUM IMPREGNATED AMBERSORB 572 (1500 ppmy TCE feed).

CATALYST	Temperature (°C)	EBCT (s)	Conversion (%)	Mineralization (%)	Water (Y/N)
-Ti	105	12.7	>99.9	90	N
	105	12.7	>99.9	77	Nª
	105	12.7	>99.9	28	N <sup>b</sup>
	150	23	>99.9	20	N
	175	35	>99.9	50	N
	175	10	>99.9	30	N
	175	3.7	>99.9	50	N
$-\mathrm{Ti}(\mathbf{W})^{\dagger}$	200	3.3	see text		
	200	31	>99.9	NT	N
	200	11	<99.9	58	N
	200	11	>99.9	35	N
	200	11	>99.9	20	N
	200	3.0	50	36	N

NT - Not Taken

Tabled data are a subset of 320 samples

An experimental catalyst, XE-706, based on the same technology presented here but using a proprietary metal oxide mix on Ambersorb 572, was provided by the Rohm and Haas Company. At 260 °C, with a dry 1500 ppmv TCE feed, a >99.9 percent

<sup>†</sup> Washed - see text.

<sup>&</sup>lt;sup>a</sup> After 1300 minutes.

<sup>&</sup>lt;sup>b</sup> After 1600 minutes.

<sup>&</sup>lt;sup>c</sup> 16 hours later.

<sup>&</sup>lt;sup>d</sup> 30 hours later.

conversion at an EBCT of 11 s was observed after an induction period of approximately 2 days. Small amounts of chloroform and PCE were noted in the effluent. When the feed stream was humidified the catalyst rapidly, but not completely, deactivated (although it was still losing activity when the water was removed) resulting in a TCE effluent concentration of about 400 ppmv. Within 12 hours after removing the water the catalyst had regained the >99.9 percent conversion level before water was added. No chloride analyses were done to confirm mineralization.

# 2.3.4 Electric Field Affects Catalytic Oxidation on MTU-CAT

Electric fields were observed to alter the by-product spectrum of TCE oxidation over MTU-CAT. The conditions of the experiments are listed in Table 2-8 and are interspersed with the experiments shown in Table 2-3. That is, thermal-only operations were performed (heat tape designation in Table 2-8) and then the Joule heating work was performed at the same temperature and flow conditions prior to changing to the new temperature and flow conditions were evaluated.

As discussed in section 2.3.2, the 1500 ppmv TCE dry feed (EBCT varied from 43 to 2.9 s) was converted by over 99.9 percent to mineralization products (HCl and  $\text{Cl}_2$ ) and a small amount of carbon tetrachloride (CCl<sub>4</sub>) at 200 °C. The heat-tape, used to initially heat the bed, was turned off and Joule heating maintained the bed temperature. An application of 2.5 V DC across the annular reactor containing 8.8 gm of catalyst caused the small CCl<sub>4</sub> peak to disappear from the effluent and no new peaks appeared. The cell potential was increased to 6.0 V DC to maintain the reactor at 200  $\pm$  10 °C. Despite the continued >99.9 percent conversion, the chloride results showed an 80 percent decrease in the mineralization occurring in the system with the applied electric field. Changing back to heat-tape and turning off the electric field resulted in two things: 1) the mineralization levels, as determined from the chloride mass balance, returned to >99 percent, and 2) the CCl<sub>4</sub> peak returned to the effluent.

A repeat of the experiment on the following day did not show this change in the mineralization as the chloride levels remained high under the 3-7 V DC applied potential, but the CCl<sub>4</sub> disappeared as before. Increasing the potential further to 9.0 V DC brought the CCl<sub>4</sub> peak back. Reducing the potential to 7.0 V DC caused the CCl<sub>4</sub> peak to disappear again. The zero-field mineralization levels had dropped overnight to 80 percent, which remained stable under the applied voltages. Alternating current (60 Hz), used both days, also showed the same trends as the DC.

TABLE 2-8. TEST CONDITIONS DURING ELECTROCATALYSIS RUNS (1500 ppmy TCE feed).

ppinv i	CE feed).			
Heat Source	Temp (°C)	Cl (%)	CCl <sub>4</sub>	EBCT (sec)
Heat Tape	200	>99	Yes	43
2.5-4 V DC	199-204	16	No	43
4-5 V DC	198-205	NT	No	43
5-5.5 V DC	201-207	5	No	43
5.5-6 V DC(R) <sup>†</sup>	199-206	NT	No	43
6 V AC	194-197	NT	No	43
10.5-11 V AC	198-210	5	No	43
Heat Tape	200	NT	Yes	43
Heat Tape	199	78	Yes	7.5
1.5-4.5 V DC	199-200	NT	‡	7.5
4.5-7 V DC	196-202	76	No	7.5
9-10.5 V DC	192-210	NT	Yes	7.5
7 V DC	202	NT	No	7.5
Heat Tape	200	76	Yes	2.9
2.7-5.2 V DC	197-204	NT	Yes°	2.9
5.2-7 V DC	195-205	· 67	Yes°	2.9
Heat Tape	183	>99	Yes	27

2.7-4 V DC	177-190	NT	No	27
4 V DC	184-186	40	No	27
4 V DC	183-185	NT	No	9.8
4 V DC	182-183	NT	No	4.8
4 V DC	180-182	59	No	3
Heat Tape	179-185	NT	Yes	3
Heat Tape	149	NT	Yes	2.9
2.7-16.5 V DC	141-159	NT	Yes	2.9
7.5-10.5 V DC	153-161	59	Yes	2.9

<sup>†</sup> Reversed polarity on cell.

The trend repeats itself when the experiments were run at 180 °C; complete conversion (>99.9), but mineralization drops by 74 percent with an applied electric field. In addition, the CCl<sub>4</sub> peak disappears with the applied potential, unless very high flow rates (0.5 s EBCT) are utilized and then it is only reduced in size. At 140 °C the catalyst remained active and had complete conversion, but mineralization under the applied potential was only 59 percent.

The effect of the electric field on the CCl<sub>4</sub> is dependent on flow rate. At the highest flow rate (EBCT 2.9s), under zero-field conditions, the TCE effluent concentration remained below detectable limits (>99.9 percent conversion) but the small CCl<sub>4</sub> concentration approximately doubled based on the areas from the gas chromatograph. Applying a potential of 2.7-7 V DC only decreased the CCl<sub>4</sub> concentration, not eliminating it entirely while the TCE concetration remained below detection limits.

<sup>‡</sup> Initially ves, but went away.

<sup>□</sup> Initial switch to AC resulted in a temperature rise to 231 °C in less than 1 minute. Temperature was back to 209 °C within 4 minutes.

o CCl<sub>4</sub> concentration reduced by ½ based on GC areas.

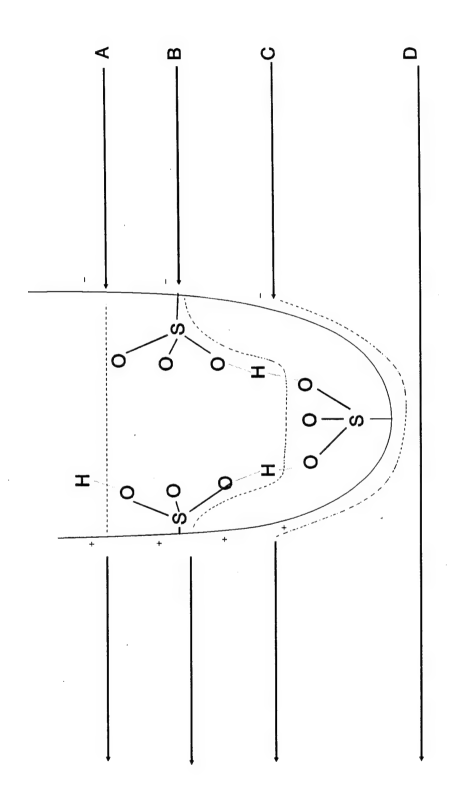
Tabled data are a subset of 192 samples.

The changes in the CCl<sub>4</sub> and chloride ion concentrations over these experiments is most perplexing. No definitive conclusion can be made based on the limited data. Uneven heating, causing hot and cold spots in the catalyst material may be one reason, another may be the how the electrons flow in the MTU-CAT catalyst.

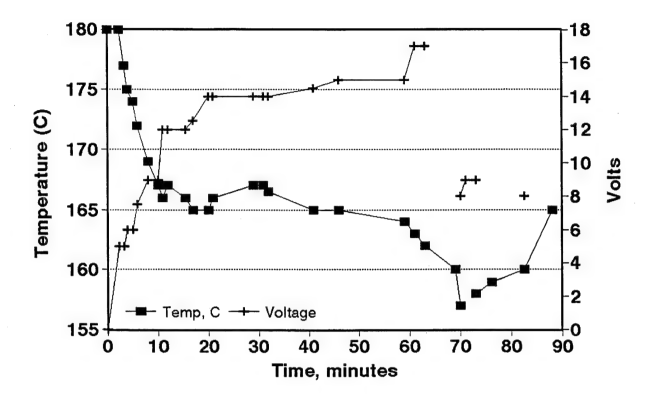
The structure of the electrically modified Ambersorb 572, MTU-CAT, may suggest how an applied electric field alters the by-product spectrum of the TCE oxidation reaction. The catalyst still contains sulfonate groups and a hypothetical catalytic site in an electric field is shown in Figure 2-7. The electron flow path differs from the bulk by traveling one of three options: 1) around the micropore (path "A"); 2) through the micropore via the sulfonic groups (path "B"); or 3) under the micropore (path "C"). The path traveled will, of course, be determined by the resistance of the different choices. If the path is through the sulfonic groups, then the electron may pass from one polymer section to another as the sulfonic groups are probably on different polymer sections. The micropore is a block to the flow of electrons and a slight electrostatic charge may build up on the sides. The potential drop across one micropore due to the overall applied electric field would be insignificant (at a bed potential of 5 V; assuming 40 beads between electrodes, no loss at electrode:bed interface with a linear drop across the beads, and a pore diameter of 10 nm would give a potential difference of 3 x 10<sup>-6</sup> V).

# 2.3.5 Electric Field Affects Catalytic Oxidation on AM572-P

Using heat tape to heat the plain adsorbent/electrocatalyst, AM 572-P, to 180 °C in the annular reactor resulted in complete conversion of the 1500 ppmv TCE feed, while mineralization was approximately 70 percent, with CCl<sub>4</sub> just above detection limits as the only peak in the effluent. Applying an electric potential of 5-17 V DC had no effect on conversion. In direct contrast to the MTU-CAT performance, the CCl<sub>4</sub> desorbed from the surface increasing its effluent concentration by a factor of about 6 at the peak of the desorption (see Figure 2-8). While the external thermocouple registered a lower temperature, the resistance change in the bed indicates an increased temperature in the



Electron paths: "A" behind pore; "B" through sulfonic groups; "C" under pore; and "D" bulk flow. Figure 2-7. Hypothetical Micropore in an Electric Field.



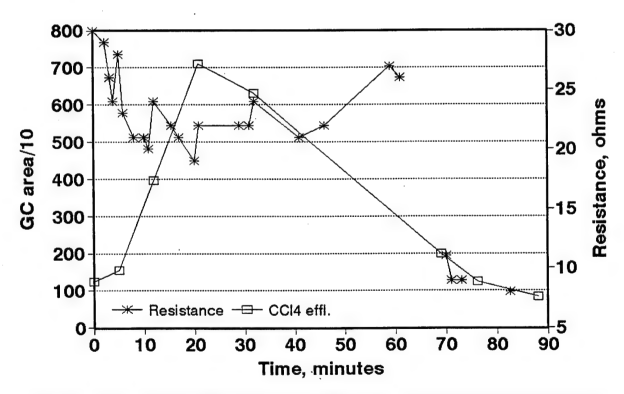


Figure 2-8. CCl4 Effluent over AM572-P under Dynamic Electric Field in Annular Reactor; EBCT = 16.7s, 1 atm (no heat tape)

39

bed, suggesting a thermal gradient was set up in the reactor. Unfortunately this means the true temperature at the adsorbent surface is unknown.

Ambersorb 572 was impregnated with a series of transition metal oxides (titanium, cobalt, and manganese) to create catalytic oxidation sites inside the polymer matrix as described earlier. The titanium oxide catalyst was initially non-conductive but became conductive upon washing with Milli-Q water. An applied electric field over these catalysts in the same annular reactor did not alter the performance of the thermal-only catalysis. Cation migration of the metals in the bed is possible though. After the cobalt oxide electrolytic experiment the center graphite rod had a bluish coating that was assumed to be cobalt.

#### 2.4 CONCLUSIONS

Low temperature catalysis of TCE over polymeric adsorbents is possible, even down to 110 °C in dry air. Water, and possibly methane, fouls the MTU-CAT catalyst, but water has no effect on the plain Ambersorb 572 catalyst. The fouling that does occur is reversible. Impregnation of metal oxides do not enhance the mineralization or conversion of the parent material to oxidize TCE under the conditions of this study. Sulfonic-group supported carbenium ions or radicals potentially exist and are probably the catalyst responsible for the low temperature conversion of TCE. By-products vary slightly and are chloroform, carbon tetrachloride, perchloroethylene, hexachloroethane, and, under extremely dry conditions, phosgene.

While this work points to the possible operation of this catalyst at temperatures of 110 °C, it should be realized that there are extensive by-products formed and complete mineralization is not realized. In addition, the operation time of the catalyst is severely shortened as the temperature is decreased as indicated by the significant increase in by-products detected by the gas chromatograph. However, if the suspected radical/carbenium ions are present and acting as the reactive intermediate species, then

increasing the temperature too far may either destabilize the -SO<sub>3</sub>H system or simply desorb the oxidizing intermediates.

An imposed electric field can minimize or eliminate, depending on flow and potential, carbon tetrachloride (a by-product from the catalytic oxidation of TCE) over an Ambersorb 572-based catalyst. At high flow conditions the by-product is merely reduced and not eliminated. Impregnation of Ambersorb 572 by transition metal catalysts did not demonstrate any impact from an applied field. At potentials above 7 V DC the effect is reversed and carbon tetrachloride returns to the effluent.

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Ambersorb® is a trademark of the Rohm and Haas Company, Philadelphia, PA. Milli-Q® is a trademark of the Millipore Company, Molsheim, France.

# Chapter 3

Adsorption and Destruction of Trichloroethylene on Carbonaceous Adsorbents Under Applied Electric Fields

#### 3.1 INTRODUCTION

Technologies used to control off-gas emissions are typically adsorption onto activated carbon or oxidation, either in a high temperature (>2000 °C) flame combustor or at lower temperatures (200-500 °C) over a catalyst. Adsorption capacities of carbon can be severely limited due to high humidities entering the carbon bed. At high water vapor pressures, that is, high relative humidities, capillary condensation occurs inside the carbon, drastically reducing the adsorption capacity of the bed. Crittenden, et al (1988) demonstrated that heating the off-gas to lower the relative humidity (RH) was cost effective to maintain the full carbon bed capacity.

Once the carbon is loaded with organics it must be regenerated, usually with steam, and the resulting liquid requires further treatment such as incineration or catalytic oxidation although there is increased interest in reuse and recovery of the organics. Fuel costs for the high temperature oxidation systems usually limit this technology to the most recalcitrant organics (e.g. PCBs). Even catalytic oxidation units operating at moderate temperatures consume energy, increasing operating costs.

Research continues to focus on energy efficient alternatives to regenerate carbon and also on catalysts (or other organic destruction processes) that operate at lower temperatures (<300 °C). The alternatives focused on here relate to using electric fields to regenerate adsorbents, to utilize a novel RH swing process that efficiently adsorbs and concentrates organics, to drive low-temperature (140-200 °C) catalytic oxidation reactions, and even to directly electrooxidize organics.

Lordgooei, (1995) and Petkovska et al, (1991) utilized electrical energy regenerate adsorbents. The process thermally heats a bed of carbon fibers while using a low-flow purge rate to remove the organics. This concept was originally used by Economy and Lin (1976) when the carbon fiber adsorbent was first introduced into the marketplace. The resistive, or Joule, heating drives the organic off the adsorbent and into the surrounding air where it is swept away by the purge gas. Joule heating occurs when the electrons flowing through the material collide with the lattice structure. The kinetic energy is converted to heat upon impact. The greater the number of electrons flowing (larger current) the more collisions occur causing the adsorbent to increase in temperature. Accordingly, increasing the irregularities in the lattice structure increases the resistance, reducing the current needed to obtain a given desorption temperature.

Petkovska et al (1991) regenerated a stack of 20 carbon cloth sections that were loaded with 1,1,1 Trichloroethane using electrical energy at a maximum temperature of 150 °C. Using direct current (3.5-5.4 V; 0.13-0.17 V/cm²) he found that a high voltage (that is, high temperature) and low purge rate are the best combination to concentrate the organics for further treatment either by condensation or destruction, which concurs with the findings of Davis (1987) for thermal treatment of adsorbents. Petkovska went on to develop a one-dimensional model (Petkovska and Mitrović, 1992) to describe the electrothermal desorption process. Unfortunately the irregularities of the material cause the bed to be non-uniformly heated, a requirement for the Petkovska model. Lordgooei (1995), cognizant of the three dimensional temperature variation in the carbon bed that develops during the heating cycle, is working on a numerical model to describe the adsorption/regeneration system. The Joule heating technology is also applicable to granular activated carbon (GAC), and others (Katsura, 1991; Ozawa, 1978; and Stankiewicz, 1990) have pursued that process.

The water vapor isotherms on the activated carbon cloth used in this study mimic granular activated carbon, resulting in capillary condensation starting around 50 percent RH (Cal, 1995). Competitive effects between water and the organics studied was also reported. He found that benzene adsorption (500 ppmv) was not affected by water until

the feed stream reached 65 percent RH, when a rapid decrease in the benzene capacity was noted with increasing RH. The water condensed in the micropores blocking the pores to the gas phase benzene. However, increased benzene concentrations decreased the amount of water adsorbed at a given RH (at 86 percent RH and 500 ppmv, 284 mg water/g adsorbent was adsorbed - at the same RH but at 1000 ppmv benzene, only 165 mg water/g adsorbent was adsorbed).

Since the cloth adsorbent behaves like a semiconductor (Petkovska et al, 1991) it would be expected that the material would behave uniquely in an electric field.

Vol'kenshtein (1963) predicted that the adsorption properties of semiconductors could be altered by an electric field that changed the Fermi level of the semiconductor. This electrical-field-induced adsorption, or electrosorption, has been reported in the literature over different materials in aqueous phases, including activated carbon, when high potentials are used (up to 25,000 V) (Woodard et al, 1986; Holder, 1970; Hoenig and Lane, 1968).

Static electric fields can also be utilized to directly alter reactions. Since gas phase electrocatalyst reactions occur (e.g., ammonia synthesis occurs over metal electrodes at high potentials [Yeh, 1968]), it was thought that trichloroethylene (TCE) may be directly oxidized over the carbon cloth. There is no literature on the electrooxidation of halogenated organics over any surface, metal or carbonaceous.

In this work electric fields were applied to adsorbent beds. Dynamic fields, causing Joule heating, were applied to adsorbents at low and moderate levels. The low level heating changed the local relative humidity reducing the competitive interaction of the water which resulted in an increased organic capacity in the carbon cloth. Isotherms on the cloth at elevated temperatures were conducted with and without applied electrical energy to determine the capacities of the various systems. At higher current densities the adsorbent was heated to the point of regeneration, or in the case of the Ambersorb 572®, to the point of catalytic oxidation of trichloroethylene. Static fields (+1000 V DC) were

applied to the carbon cloth to determine if direct electrooxidation of TCE is possible under lab and field conditions.

### 3.2 MATERIALS AND METHODS

Water for the humidity bubblers came from a Milli-Q® station (Millipore, Molsheim, France) in the lab. TCE in dry nitrogen (500 parts per million by volume [ppmv]) was purchased from Scott Specialty Gas Company (Riverside, CA). TCE in dry air (15 & 1500 ppmv) were purchased from Matheson Gas Products (Jolliet, IL). The Ambersorb 572 adsorbent (40x60 mesh, Lot No:923318) was purchased from the Rohm and Haas Company (Philadelphia, PA). The carbon cloth ("Kuractive," FT 200-15, Lot # FK 622096-1B, Kuraray Company, Bizen, Okayama, Japan) was provided, as were the graphite rods (The Carbide/Graphite Group, Inc., St. Mary's, PA). All carrier gases including air, argon, helium, and nitrogen were ultra-pure and purchased from Interstate Gas Company (Marinette, MI).

General lab equipment items are: a mass flow controller 5850 E (Brooks Instrument Division, Emerson Electric Co., Hatfield PA), a low voltage/medium amperage (0-42 V/3 A) DC source 6290A (Hewlett Packard), an Alnor Type 7000 Dew Pointer (Alnor Instrument Company, Chicago, IL) and a Fisher Scientific temperature/humidity monitor (Pittsburgh, PA) to determine relative humidities, and Variac controllers for external heating of the system. The Fisher Scientific humidity meter was calibrated using sulfuric acid solutions at known concentrations yielding constant relative humidities as listed in the CRC Handbook of Chemistry and Physics, 1980.

The lab bench contained a dual AC/DC power supply (Lab-Volt, model 194, Buck Engineering Co, Farmingdale NJ) with an output of 0-36 V at 0-5 Amps (AC frequency was 60 Hz). The HP DC source mentioned above could be operated in two modes: constant current or constant voltage. The Lab-Volt could only be operated in the constant voltage mode.

Sampling procedures to determine the influent and effluent concentrations of TCE, measurement of by-products, and chloride/sulfate production are reported in Chapter 2.

The adsorbent/catalyst materials were used in two different packed-bed configurations. All of the catalytic oxidation reactions were over Ambersorb 572 beads that were in an annular reactor as shown in Figure 3-1. The graphite rod (The Carbide/Graphite Group, Inc., St. Marys, PA; no part or lot number) in the center of the reactor was 1/8 inch in diameter and extended to within 1/2 inch of the bottom. A section of conductive, activated carbon cloth ("Kuractive," FT 200-15, Lot # FK 622096-1B) was used in the bottom of the reactor to keep the adsorbent/catalyst from falling out of the reactor. Electrical isolation was accomplished with Teflon® tape and glass tubing connectors.

The carbon cloth was used in a flow through system where the bed was sandwiched between two porous stainless steel electrodes as shown in Figure 4-2b. The flows were metered and inlet and effluent concentrations could be taken as shown in Figure 3-2. The reactors could be heated externally with heat tape, internally with Joule heating, or both. The larger reactor (1.5 inch I.D. by 6 inches long carbon steel column) was used in the annular configuration with thermocouples inside and outside the reactor. The flow-through reactor (quartz glass tube 1/4 inch OD; 1/8 inch ID) only had a thermocouple on the outside of the reactor.

The isotherms were conducted by passing contaminated air at a given flow rate over the adsorbent while monitoring the effluent. When the effluent reached the influent and stabilized for 10 or more hours, the feed was changed to the next higher concentration. The process was repeated until all three concentrations (15,500, and 1500 ppmv) of TCE had been fed through the adsorbent bed. The area under each of the three breakthrough curves was calculated and summed up to give the reported capacities.

The reactor beds for the static electric field experiments were made out of the same glass and screen materials as the flow through cells noted above. The field unit consisted

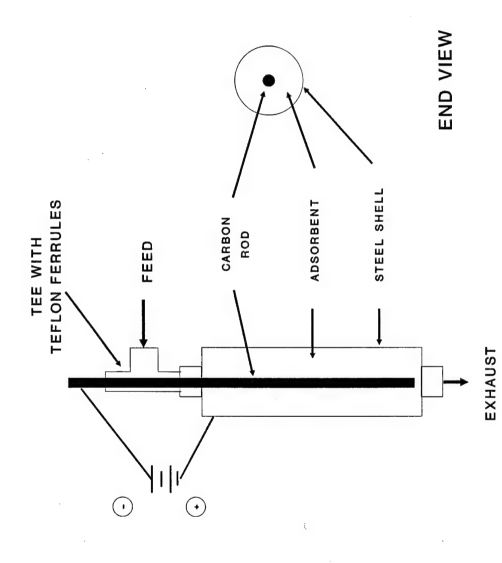


Figure 3-1. Annular Reactor Design

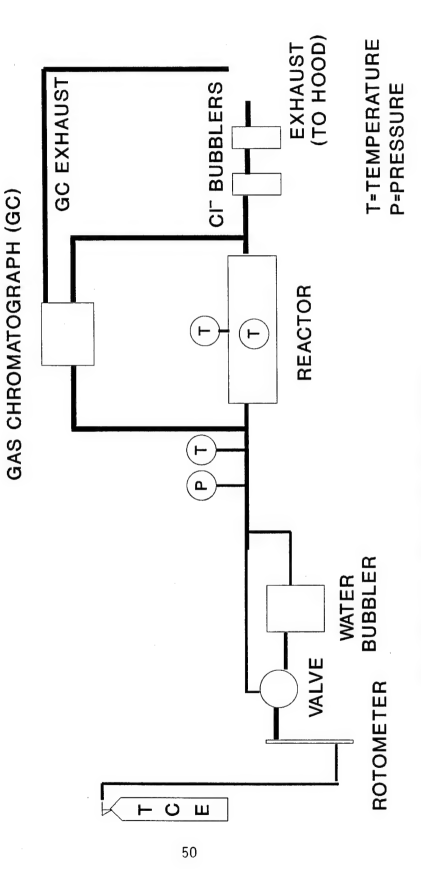


Figure 3-2. Experimental Setup

of a support/electrode stainless steel screen, with two layers of carbon cloth setting upon it with no second screen (unit was vertical with gas flowing from below, past the screen and the cloth sections before exiting). The lab reactor was a flow cell with both stainless steel screens, but the second electrode was not connected.

All the gas flow rates reported herein are based on 1 atm and 22 °C unless otherwise stated. The empty bed contact times (EBCT = volume of catalyst bed/volumetric flow rate) are based on the flow rate at 22 °C. The pressure drop across the bed at the different flowrates used in this study changed by less than 0.5 psi and did not significantly alter the vapor phase concentration.

The term conversion is determined by the amount of TCE in the exit stream as compared to the amount in the feed stream. A 100 percent conversion is defined as the complete loss of the TCE in the effluent. Mineralization is defined as the amount of HCl or Cl<sub>2</sub> found in the traps as compared to that which is expected from stoichiometry. A 100 percent mineralization level indicates that all of the available incoming chlorine is accounted for in the traps.

#### 3.2.1 Adsorbent Characteristics

The plain Ambersorb 572 (AM 572-P) is a sulfonated polystyrene divinylbenzene cross-linked polymeric adsorbent (ion exchange resin) that has been pyrolyzed in a patented process. This adsorbent comes in the form of uniform spherical beads with the characteristics shown in Table 3-1. The material contains micropores, mesopores, and macropores. Inside the pores the sulfonate groups are attached to the polymer material, probably attached as functional groups off the ends of the planar structures shown by Neely and Isacoff (1982). In the micropores, and possibly the mesopores, the sulfate groups are in close enough proximity to interact with one another to hydrogen bond and create strong acidic sites (Gates, 1992).

The carbon cloth (K-cloth), also listed in Table 3-1 for comparison, is essentially phenolic (Novoloid) fibers in a needled felt form that have been pyrolyzed and activated at high temperatures. Thus, heating the cloth to high temperatures to regenerate the adsorbent is not a problem. Unfortunately the AM 572-P is limited to 325 °C in air or breakdown of the adsorbent begins to occur. This limits the operational range of the Joule heating process for the adsorbent.

The thermal conductivity of the AM 572-P is poor and thermal gradients can develop in the beds. Using the AM 572-P in the annular reactor with thermocouples inside the bed and on the outside edge, thermal gradients of 10 degrees Celsius per centimeter (10 °C/cm) were observed during initial heating with heat tape. This points to the use of short electrode spacing in the final reactor design. The carbon cloth has a thermal conductivity of 0.7 mW/cm °K (Hayes, 1993), approximately that of platinum (Robinson, 1987).

TABLE 3-1. ADSORBENT CHARACTERISTICS

ADSORBENT	SURFACE AREA m²/g	DENSITY g/cc	PORE VOLUME ml/g
Ambersorb 572 †	1100	0.49	0.84
Kuraray FT 200-15 ++	730-900	1.27	0.38

<sup>&</sup>lt;sup>†</sup> Data from Rohm and Haas (1992).

#### 3.2.2 Field Site Description

The field site (IC-1) is at McClellan Air Force Base (McAFB), an active installation located north of Sacramento in the Central Valley of California. Soil vapor extraction is in use at the site to remove TCE, tetrachloroethylene (PCE), and Freon 113 contamination in the vadose zone. The contaminated vapors are treated in a fluidized-bed catalytic oxidation unit utilizing a chromia oxide catalyst operating at 1100 °F and 630 cfm (Empty Bed Contact Time [EBCT]= 0.8 s). The effluent air goes to a caustic scrubber to remove any HCl or HF that may have formed in the reactor. During the field

<sup>&</sup>lt;sup>††</sup> Data from Cal (1995) and Kuraray Chemical Co.

study the incoming air concentrations varied but were typically around 15 ppmv TCE and 55 ppmv PCE (Freon 113 was undetectable on the field gas chromatograph [GC]). The feed stream relative humidity also varied from 35 to 60 percent. In the field a Scentoscreen portable GC (Sentex Systems, Inc., Ridgefield, New Jersey) was used that had an argon ionization detector in it. A mixed gas tank of TCE (100 ppmv), PCE (200 ppmv), and Freon 113 (50 ppmv) in dry air was purchased from Matheson Gas (Jolliet, IL) to mimic field site conditions. This standard gas was used in the field to calibrate the GC on-site.

### 3.3 RESULTS AND DISCUSSION

### 3.3.1 Carbon Cloth Capacities for TCE

Isotherms were performed on the cloth to evaluate the impact of applied electrical potential on the adsorption capacity and relative humidity. Figure 3-3 shows the TCE (15, 500 and 1500 ppmv) isotherm results at 38 °C with dry and humid feeds; with and without electric potential. Heat tape, controlled by a temperature controller, was used to keep the bed at isothermal conditions. As the amount of Joule heating increased, the heat input from the heat tape automatically reduced, maintaining the temperature of the bed at 38 °C (± 0.3 °C). As shown in Figure 3-3, no increase in capacity was noted with an applied potential. Indeed, there appears to be a negative influence, however the difference is within experimental error. This indicates that the effect at these potentials is due to the changes in the relative humidity and not due to electrosorption occurring on the surface. As can be seen, the capacity is reduced at the 1500 ppmv level with water addition, regardless whether the electrical potential is there or not.

Humidity (80 percent RH) has a much larger impact on the capacity than temperature. A 15 ppmv TCE dry feed at 25 °C (not shown in Figure 3-3) resulted in essentially the same capacity as the dry 38 °C loading. However, the difference between the dry and humid 25 °C loadings (a factor of about 9) is due to the loss of capacity from relative humidity effects. Localized Joule heating of the bed decreases the RH and

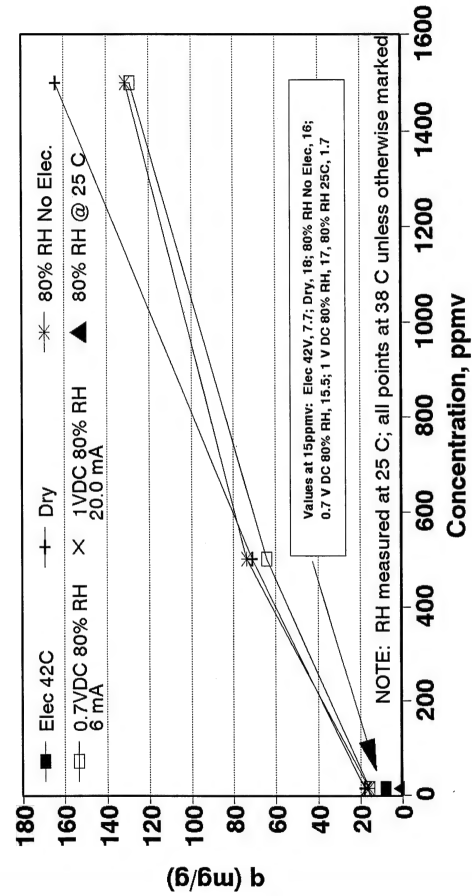


Figure 3-3. Cloth Isotherms at 38 C in Trichloroethylene Flow Cell

dramatically increases the capacity of the bed over the non-heated capacity. However, heating the bed too much results in a decreased adsorption capacity (see Figure 3-3; 42 °C, 15 ppmv, isotherm point)

# 3.3.2 Joule Heating

When discussing Joule heating it is important to address the type of electrical energy used (AC or DC), the magnitude of the applied potential, and the reactor design (packed bed or annular). Another factor is the distance between electrodes (length of resistor). Increased resistance is desirable to keep the power requirements low for a given heating cycle, however hot and cold spots can develop in a large bed with large distances between electrodes.

# 3.3.3 Joule Heating of Activated Carbon Cloth

Low level Joule heating of the carbon cloth in humid feeds showed an increased capacity with the application of a small potential over the initial room-temperature capacity. Figure 3-4 illustrates this effect. The initial short breakthrough is at 25 °C and 80 percent RH. An applied potential of 4 VDC (30 mA) raised the bed temperature to 42 °C and caused the effluent to drop, an indication that the bed was adsorbing more TCE. At an EBCT of 0.1 s the exiting air temperature was only 0.5 to 1 °C warmer than the inlet air temperature. The increase in capacity is approximately 4.5 times (the capacities are also shown in Figure 3-3 as the humid feed at 25 °C and electroheated capacity at 42 °C). This is not an optimized system and capacities greater than this are expected. Based on the data in Figure 3-3 at 38 °C the capacity increase could be as much as 9 times greater than the non-heated capacity. This increase in capacity is due to the competing effects within the adsorbent. As the adsorbent gets hotter, the overall capacity is reduced due to thermal interactions, however the relative humidity is also reduced and the capacity for the organic increases.

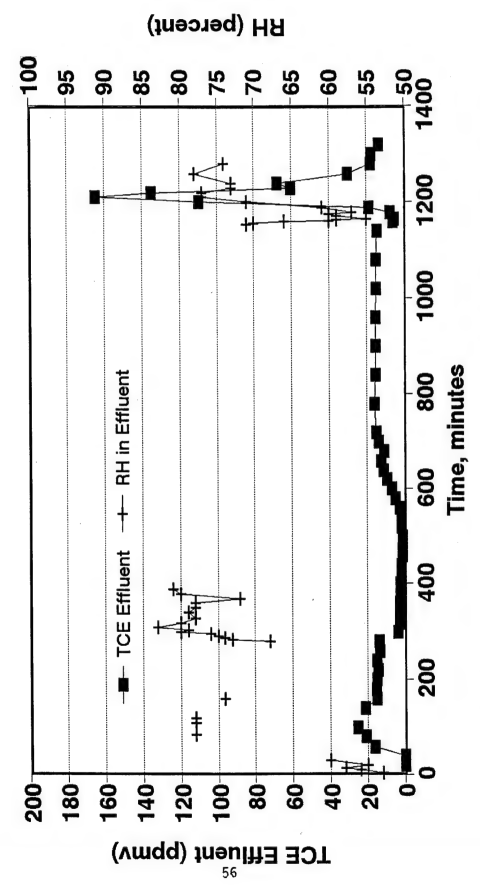


Figure 3-4. Humidity/Joule Effect on Carbon Cloth, Humid 15 ppmv TCE Feed, 0.1 s EBCT, 1 atm

When the applied potential is removed the excess adsorbed TCE is rapidly desorbed with a peak maximum concentration of 160 ppmv, or about a ten fold increase in concentration. The cycle is repeatable, with the same capacity and desorption curves resulting. The power input to create this additional capacity is 0.6 w-hr (36 w-hr/gram adsorbent). An alternative process (Crittenden et al., 1988) would heat the incoming air to reduce the humidity, instead of locally reducing it at the adsorbent surface. The incoming air has an enthalpy at saturation of 36.3 Btu/lb dry air. At 42 °C the RH would have dropped to 33 percent and the enthalpy at saturation would be 45.3 Btu/lb dry air. Heating to 42 °C the same volume of air that passed over the adsorbent during the increased capacity time would have required 0.1 w-hr of power, or 1/6th the amount from the Joule heating. The system is not optimized as to design or materials and the factor of six difference may be a conservative estimate.

The heating input to raise the bed to 42 °C is not factored in the above numbers for the air-heating option, but is estimated to be 0.05 cal, based on a Specific Heat of carbon at 0.168 cal/g-°C (Robinson, 1987). Based on the small thermal increase of the incoming air over the Joule heated bed, it is expected that the heat transfer to the bed would be slow and it would take some time to heat the bed initially. Therefore, an optimized system would utilize Joule heating to rapidly heat the bed initially to the desired temperature and then utilize incoming air that was preheated to maintain the desired bed temperature to reduce the RH over the adsorbent.

### 3.3.4 Joule Heating of Ambersorb 572

The electron flow within the bed of Ambersorb beads is expected to flow radially through a given cross section of the bed. Due to the design of the reactor the actual electron density will be higher at the center graphite rod, however. Inside the individual particles variations in current density are also probable. Due to the porous nature and physical makeup of the polymer, the electron flow inside the particle may take a tortuous path within the bead, traveling from one pyrolized polymer section to another, possibly even by way of the sulfonate groups (if those groups are in close enough

proximity to allow electron transfer). If the adsorbent is heated too rapidly these electron flow paths may heat portions of the bead beyond safe limits, even though the bulk bed temperature is well below the temperature limit of the material. Indeed, MTU-CAT, a Joule-heated catalyst created from plain Ambersorb 572, was heated rapidly changing the pore shape and catalytic performance as discussed in Chapter 2.

Joule heating, at sufficiently high potentials, can provide the thermal energy of activation for the catalytic oxidation of TCE. Plain Ambersorb 572 has been shown to have catalytic properties (Chapter 2, this work; Petrosius et al, 1993; and Petrosius and Drago, 1992) for the oxidation of chlorinated organics.

As discussed in Chapter 2, heat tape was used on an annular reactor to provide the initial 200 °C heating input to drive the oxidation reaction with TCE. This reduced the incoming, dry, 1500 ppmv feed, to less than 1 ppmv (detection limit on gas chromatograph). Joule heating, both AC (60 Hz) and DC, was used to replace the heat tape and the TCE effluent concentration remained below detection limits over multiple testing sequences. Reducing the temperature of the reaction to 180 °C and repeating the multiple testing sequences gave the same result: complete conversion of TCE in the reactor. A limited number of experiments were done even as low as 140 °C, with no detectable TCE in the effluent. A summary of the conditions used are shown in Table 3-2.

#### 3.3.5 Electric Field Influences on Regeneration

Regeneration of adsorbents can be done with Joule heating alone, or in combination with steam. Hand, et al (1995) showed effective regeneration by using a combination of electricity and steam to clean samples of Ambersorb 572 which had been loaded with TCE in the liquid phase. The steam-alone process had broader peak effluents and longer tails, while the electroenhanced process had higher, sharper effluent profiles and almost no tailing. This is due to the internal heating of the bed by the Joule heat, increasing the internal thermal energy of the bed. This may have kept the steam from condensing in the pores and slowing mass transfer out of the adsorbent.

TABLE 3-2. TEST CONDITIONS DURING JOULE HEATING EXPERIMENTS (1500 ppmv TCE feed).

Voltage Applied	Temperature (°C)	EBCT (sec)	
2.5-4 V DC	199-204	43	
4-5 V DC	198-205	43	
5-5.5 V DC	201-207	43	
5.5-6 V DC(R) <sup>†</sup>	199-206 43		
6 V AC	194-197	43	
10.5-11 V AC	198-210	43	
1.5-4.5 V DC	199-200	7.5	
4.5-7 V DC	196-202	7.5	
9-10.5 V DC	192-210	7.5	
7 V DC	202	7.5	
6-10 V AC	198-209°	7.5	
2.7-5.2 V DC	197-204	2.9	
5.2-7 V DC	195-205	2.9	
2.7-4 V DC	177-190	27	
4 V DC	184-186	27	
4 V DC	183-185	9.8	
4 V DC	182-183	4.8	
4 V DC	180-182	3	
2.7-16.5 V DC	141-159	2.9	
7.5-10.5 V DC	153-161	2.9	

<sup>†</sup> Reversed polarity on cell.

In regeneration experiments with the liquid-loaded AM 572-P using Joule heating alone, the process did not work very well. As the adsorbent dried it clumped

Initial switch to AC resulted in a temperature rise to 231 °C in less than 1 minute. Temperature was back to 209 °C within 4 minutes.

together, pulling itself away from the electrodes in the annular reactor. This loss of electrical contact caused problems of uneven heating. This problem could be avoided by switching to a different reactor design, using different materials, or both. For example, if the reactor was agitated by an air blast to break up the clumps, this would allow the current to flow again. The liquid-loading work of Economy and Lin (1976) with phenol showed that the cloth fibers do not show this tendency and could be effectively regenerated from liquid loadings.

There is a more important drawback to regenerating these adsorbents with Joule heating however. The resistance of the carbon cloth decreases when the material is heated (the semiconductor behavior noted by Petkovska, 1991). As the adsorbent heats, the hot areas become more conductive and more current flows through that section of the bed. Of course, as more current flows, the hotter that section becomes and the process is in a vicious cycle until the adsorbent is red hot and glowing. Indeed, in one early experiment a copper electrode end was melted (mp 1083 °C) during a rapid heating cycle. In addition, as discussed in the next chapter, when an organic adsorbs to the cloth the resistance increases. Thus, the areas that need the regeneration the most are also the most resistive and the electron path will favor "clean" adsorbent sections since they are more conductive.

Care can be used to minimize this effect by proper electrode configuration, spacing, and heating rates. For example annular heaters can be used to evenly distribute the charge along the length of the bed, obtaining a radial direction of electron flow and heating. However, the adsorbent in the interior would experience a higher current density and a thermal gradient in the radial direction would develop. A better alternative would be to have large flat plates or flow through screens to get an even distribution of current prior to going through the adsorbent bed. In preliminary experiments it is anticipated that the electrical costs for regeneration are 16 \$/m³, based on a 200 °C heating cycle for 10 hours. This is in comparison to 110 \$/m³ for steam regeneration (assuming 20 Kg steam/Kg adsorbent, and \$5/1000 lb steam [Rick et al, 1987])

## 3.3.6 Electrostatic Field Affects over Carbon Cloth

Laboratory and field experiments were performed to evaluate the impact of Joule heating on the catalytic oxidation of TCE over carbonaceous adsorbents. The static fields are applied at ambient temperatures, however, some current flows during the experiments as indicated by the small increase in bed temperature. The laboratory feed consisted of just trichloroethylene (TCE), whereas the field feed stream has TCE, tetrachloroethylene (PCE), and Freon 113.

. Using 4 sections (0.007 g) of the carbon cloth in a 1/4" OD flow-through reactor, the effects of a static electric field were first investigated in the laboratory. A stream of humidified TCE in air (15 ppmv; EBCT = 0.1 s) was passed through the bed. Sufficient volume was used to reach capacity loading at room temperature (effluent=influent). A static potential of +1000 V was applied to the effluent-end stainless steel screen holding the cloth in position. The temperature of the bed rose 6 °C above room temperature within 20 minutes and remained stable. The effluent decreased due to the relative humidity shift described above. After 15 hours the effluent TCE concentration was about 60 percent of that coming in. In addition, there were 4 new peaks on the chromatogram output. Based on retention times three of them were tentatively identified as carbon tetrachloride, chloroform, and PCE. The fourth peak was an unknown. Unfortunately, when attempting to get a chloride sample the reactor broke losing any other confirmatory information on the reaction. Time constraints for the field work did not allow for a repeat experiment.

Direct electrooxidation of TCE was attempted again under field conditions at McAFB, Site IC-1. Two sections of the carbon cloth were supported on a stainless steel screen that was charged to +1000 V DC. The contaminant stream varied from 35-55 percent relative humidity and flowed up through the screen, through the bed and to chloride traps (two in series). No heat control was used on the bed, and ambient temperatures ranged from 17-38 °C. Field results showed no difference between influent and effluent TCE/PCE concentrations. Chloride analysis of the traps verified this, but also

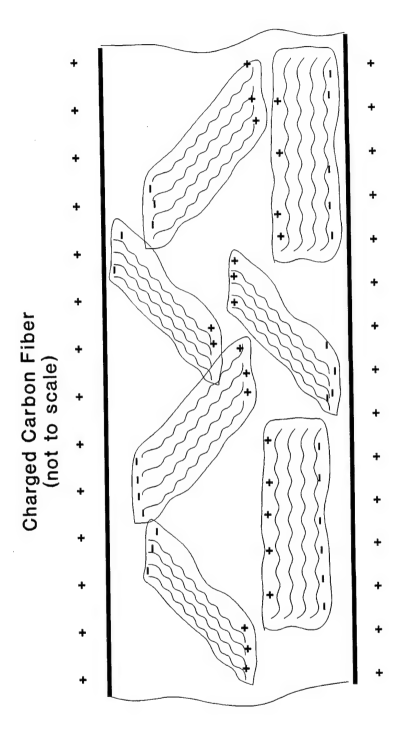
showed fluoride in the effluent. Thus, while TCE and PCE were not being affected, the Freon 113 was at least partially oxidized.

Under these electrostatic conditions the carbon cloth individual fibers would have been charged on the outer surfaces of each rod-shaped fiber. Some electron flow did occur, based on the Joule heating that was noted. The barriers between two graphitic-ordered domains, separated by an amorphous sea, would have caused induction of the charge as shown in Figure 3-5. These charge separations at the boundaries may have had some impact on the reaction.

Under electrodynamic conditions the charges would have been spread throughout the rod not just on the outer shell of the fiber. In order to take advantage of the high surface area for catalysis (and the sulfonated groups in the Ambersorb-based catalysts) electrodynamic conditions need to be used. However, when this is done, there is Joule heating that also occurs and separating the thermal vs. electrolytic effects are not possible.

### 3.4 CONCLUSIONS

Joule heating can rapidly heat the adsorbents used in this study as well as enhance the desorption rate during steam regeneration. However, hot spots can occur in an adsorbent bed and controls (bed design, electrode spacing, heating rate) are critical to limiting their effects. Regenerating liquid loaded Ambersorb 572 with Joule heating is problematic and a final design will require special agitation tools. Low level Joule heating can increase the capacity of the carbon cloth for TCE in humid streams. However, once the bed is hot, the energy costs indicate that heating the incoming air would be cheaper than heating the bed with Joule heating. In addition, cooling the bed back down in a humid stream removes TCE from the surface, not completely regenerating it but rather releasing the excess adsorbed under reduced RH conditions. Thus an adsorption/regeneration cycle could be used for just the increased capacity of the adsorbent due to the RH shift to concentrate the organics in the air stream. In strong electrostatic fields (+1000 V DC) direct



Charge in Crystallites Within Carbon Fibers Figure 3-5. Electrostatic Charge on Fiber Induces

electrooxidation of TCE in the lab, and Freon 113 in the field, is suggested based on the limited data. Further work is needed to verify the finding.

#### 3.5 ACKNOWLEDGEMENTS

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Ambersorb® is a trademark of the Rohm and Haas Company.

# Chapter 4

Electronic Factors of Adsorption on Carbon Fibers

### 4.1 INTRODUCTION

Gas phase adsorption occurs on the surface and in the pores of the adsorbent. In physisorption the adsorbate (the substance being adsorbed) is held by a combination of van der Waals and electrostatic forces. In contrast, chemisorption occurs with the actual transfer of electron(s) between the adsorbent and the adsorbate. The adsorption phenomenon is usually categorized as physisorption and chemisorption (Sontheimer et al 1988) and it is difficult to distinguish them at times. In micropores the forces from the adjacent walls overlap creating high energy adsorption sites. Chemisorption at these sites may occur while physisorption is simultaneously occurring at lower energy sites, including surface sites not associated with pores. In addition, different molecules with different dipole moments and sizes will have different affinities for various adsorbents.

Two forms of activated carbon cloths (ACC) made from fibers were used as adsorbents in this study. The first form, predominantly used in this study, is a non-woven material with the consistency, texture, and appearance of felt. The second form is a cloth weave where the fibers are spun into long strands which are then woven into a cross pattern to yield a cloth-like appearance. Hayes (1994), in company literature from American Kynol, gives an excellent treatise on how the fibers are made from phenolic precursors (Novoloid fibers). He describes the pyrolyzation process and ways to modify the surface of the materials for a variety of applications. Some characteristics of the fibers are listed in Table 4-1. The impact of an applied potential on the adsorption of organic compounds was evaluated. The organic compounds are of industrial importance and span a wide range of adsorption strengths.

TABLE 4-1. CHARACTERISTICS OF ACTIVATED CARBON MATERIALS

	Cloth Weave Needled Felt		
Diameter, $\mu$ m	10	10	
Density, g/cc	1.4-1.6	0.1	
Surface Area, m <sup>2</sup> /g	1500	1500	
Thermal Conductivity,		0.72	
W/m°K			

From Hayes (1993, and 1994) and Kuraray Company, Inc.

The carbon fibers are pyrolyzed and activated in one step (Hayes, 1994). The resulting fiber is approximately 95 percent carbon and contains crystallites that are sets of graphitic, or basal, planes stacked one upon another as shown in Figure 4-1. The area between the crystallites is amorphous carbon, probably tetrahedrally bonded, and other impurities (e.g. oxygen, water, etc.) may also be in these interstitial areas. These zones between the crystallites are prone to oxidation during the activation process as the graphitic planes are quite stable. Thus major portions of the areas between the crystallites are etched away during the activation process as shown by the shaded area in Figure 4-1. If the activation process continues then the edges of the crystallites are also etched away. These opened areas become the micropores for adsorption. The crystallites are in random order, giving the fiber an overall amorphous appearance when examined spectroscopically, such as X-ray diffraction reported by Hayes (1994).

Intra particle electron conduction in the crystallites is favored along paths that are parallel to the basal planes as shown by path "A" in Figure 4-1. Conduction perpendicular to the basal planes as shown by path "B" in Figure 4-1 is possible, but it is not the preferential path. Inter particle, or transitions between particles, may occur by hopping as indicated by path "C" in Figure 4-1. The conductive polymer literature describes electrical conduction in polymers in similar terms with the charge carrier hopping from one crystallite to another with distances up to 7 Å (Thakur and Elman, 1989).

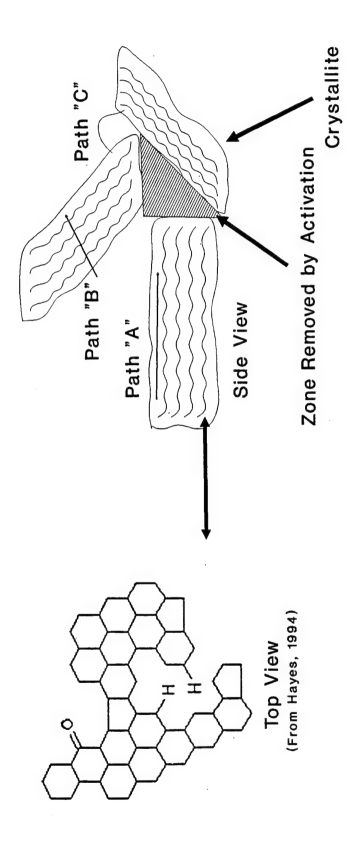


Figure 4-1. Electron Paths in Graphitic Planes and Crystallites in Cloth Fibers

The micropores in the ACC used in this study account for its large surface area. Using nitrogen adsorption data, Cal (1995) reported that all of the pores for the ACC-15 sample (another Novoloid-based activated carbon cloth made by Kynol Inc., similar to the woven structure in this study) were in the micropore region (d < 2 nm). This concurs with the report from Hayes (1994) which showed essentially no mesopores in the materials. Hayes also mentions the electrical conductivity of the cloth, and even suggests using it in capacitors because of its high surface area.

Granular activated carbon (GAC) has a different structure, containing micropores, mesopores, and macropores, however the two adsorbents, ACC and GAC, are similar in their adsorption trends. Recent work (Cal, 1995) and related on-going work (Lordgooei, 1995) at the University of Illinois at Urbana-Champaign utilizes the activated carbon cloth made from Novoloid fibers. Cal found that the carbon fibers have the same general adsorption trends as GAC. That is, compounds adsorbed strongly to GAC were found to be strongly adsorbing to the carbon fiber and visa versa. In addition, he found that water competes for the available adsorption sites.

While no references can be found relating the adsorption of organics to resistance change on carbon fibers, others have explored physical changes that can occur when using the cloth material. A group of French researchers (Baudu et al, 1992) found an empirical relationship for the resistance change the carbon cloth undergoes when heated. They modeled both carbon fibers and GAC, resulting in an equation that describes an exponential decay in the resistance with increasing temperature.

In this study seven compounds with varying adsorption potentials were tested to determine the impact of adsorption on activated carbon cloth and the resulting changes in resistance. Static (no flow) and dynamic (with flow) isotherms were conducted to determine the effect of concentration, single solute effects, and water-component interactions on the resistance change in the adsorbent.

### 4.2 MATERIALS AND METHODS

Tanks of prepared gases were used for the flow through (dynamic) isotherms. Trichloroethylene (TCE) in dry nitrogen (500 parts per million by volume [ppmv]) was purchased from Scott Specialty Gas Company (Riverside, CA). TCE in dry air (15 & 1500 ppmv) was purchased from Matheson Gas Products (Jolliet, IL). The gases were used without drying.

The carbon cloth came in two forms and both were provided by the Kuraray Chemical company (Bizen, Okayama, Japan). The non-woven, felt-like material called "Kuractive," (FT 200-15, Lot # FK 622096-1B) is used the most in this study and referred to as "activated carbon cloth". While the woven material is also called "Kuractive" (CH700-15, Lot # CKA 630042-2), it is referred to as "woven" material in this study. The carbon cloths were used "as-received" and not pre-treated prior to use.

All chemicals used in the experiments and analyses were reagent grade or better. Reagent grade dichloromethane (99.9+%), methanol, carbon tetrachloride, tetrachloroethylene, trichloroethylene, and toluene were purchased from Aldrich Chemical Company (Milwaukee, WI). The water came from a laboratory Milli-Q station (Molsheim, France).

For the dynamic isotherms on-line gas-phase samples were analyzed according to the Environmental Protection Agency's Reference Method 23 (Scott Environmental, 1983) using a Hewlett Packard Model 5880A gas chromatograph equipped with a flame ionization detector (FID). Compound separation was performed using a packed column (3 percent DBP 1500, 80/120 Carbopack® B) (Supelco, PA). The gas samples passed through stainless steel lines into a six-port valve (Model RD6P, Valco) that was heated to 160 °C (Valco Instrumentation Temperature Controller). A 50  $\mu$ L sample loop attached to the six-port valve provided a gas sample size that yielded a TCE detection limit of about 1 part per million by volume (ppmv).

The experimental set-up used for the static single solute resistance measurements is shown in Figure 4-2a. Pure phase solutions of the chemicals were injected with a disposable syringe (Syringe & PrecisionGlide Needle, 322IM1½ 23GTW, Becton Dickinson & Co., Franklin Lakes, NJ) below the cloth samples, resulting in air concentrations in the isotherm bottle equal to the vapor pressure of the organic at room temperature. Excess was used to ensure complete saturation of the cloth. Nichrome wires held the cloth in place and acted as the electrodes.

The dynamic isotherms were taken in a flow through glass cell (1/4 inch OD, 1/8 inch ID) with stainless steel screens to hold the cloth sections in place as shown in Figure 4-2b. Only trichloroethylene dynamic isotherms were conducted on stacks of circular cloth sections. All dynamic adsorption experiments used 12 cloth sections, except for the work shown in Figure 4-5 which used 48 sections. Nichrome wire was used to connect to the stainless steel screens, with the other end contacting the interior of the stainless steel lines before and after the glass tubing. Exterior connections were then made to the volt-ohm meter using hose clamps and copper wiring (12 gauge).

All experiments were conducted at room temperature and 1 atm unless otherwise noted. The pressure drop across the bed at the different flowrates used in this study changed by less than 0.5 psi and did not significantly alter vapor phase concentrations.

#### 4.3 RESULTS AND DISCUSSION

#### 4.3.1 Adsorbent Characteristics

The fibers in the adsorbent are approximately 10 microns in diameter and have various lengths. The carbon cloth, or needled felt, is a jumble of the fibers entangled together into a cloth. The cloth weave is a series of fibers (40-50) entwined to form a strand. Three strands are woven together to form one cross-hatch material in the final fabric. The adsorbents were activated and carbonized in a single step, with a reported surface area of 1500 m²/g (Hayes, 1993). At pyrolization temperatures above 700 °C the

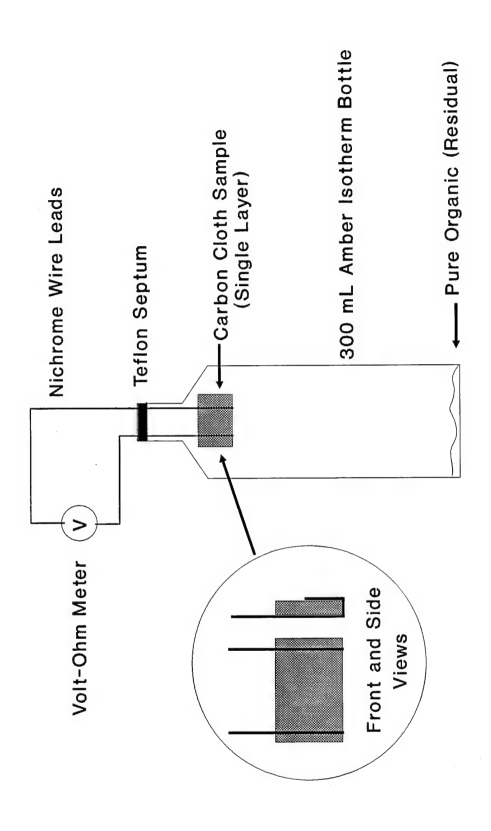


Figure 4-2a. Resistance Isotherm Setup for Cloth

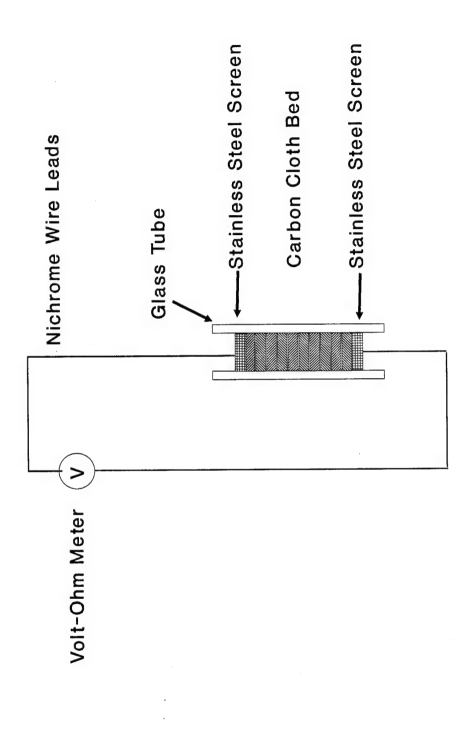


Figure 4-2b. Dynamic Resistance Isotherm Setup for Cloth

Novoloid fibers form graphitic-like structured domains with limited areas of extent that form islands in the amorphous fibers (Hayes, 1994). Fung et al (1994) report that these planes in the fibers are ~2.5 nm in diameter and occur in layers of 1-3 planes to form crystallites. Similar structures were found in carbon films where the graphitic islands are separated by regions of tetrahedrally bonded carbon (McLintock and Orr, 1968).

The carbon cloth adsorbent used in this study has the characteristic of increasing in conductivity with increasing temperature as shown in Figure 4-3. French researchers (Baudu et al, 1992) reported the following exponential relationship for the resistance changes with temperature for carbon fibers and GAC:

$$R_T = R_0 \exp[-b(1/T-1/T_0)^2]$$

where  $R_T$  is the resistance at the temperature "T",  $R_0$  is the resistance at the initial temperature " $T_0$ ", b is 98 for GAC and 83 for the fiber, and the temperatures are in degrees Celsius. Baudu et al, attribute the difference between the values of "b" to the structural differences between the different materials.

In this study these findings were not collaborated as shown in Figure 4-3. The carbon cloth was found to be linear in the temperature/resistance relationship over the temperature range used in this study. Two configurations of the cloth were used as discussed below, and also another polymer-based adsorbent, Ambersorb® 572 (Rohm and Haas, Philadelphia PA). Figure 4-3 also contains data from early work by McIntosh et al. (1947) showing a linear response with carbon rods made from carbonized zinc chloride/cellulose starting materials. It is not clear which fiber form was used in the work by Baudu et al, but the trend should remain the same for the different cloth materials.

A possible explanation for the difference was thought to be in how the measurements were taken. The cloth sections studied here were allowed to hang free in the temperature controlled oven and were not influenced by pressure due to thermal expansion differences. If the material was contained, heating could cause an increase in

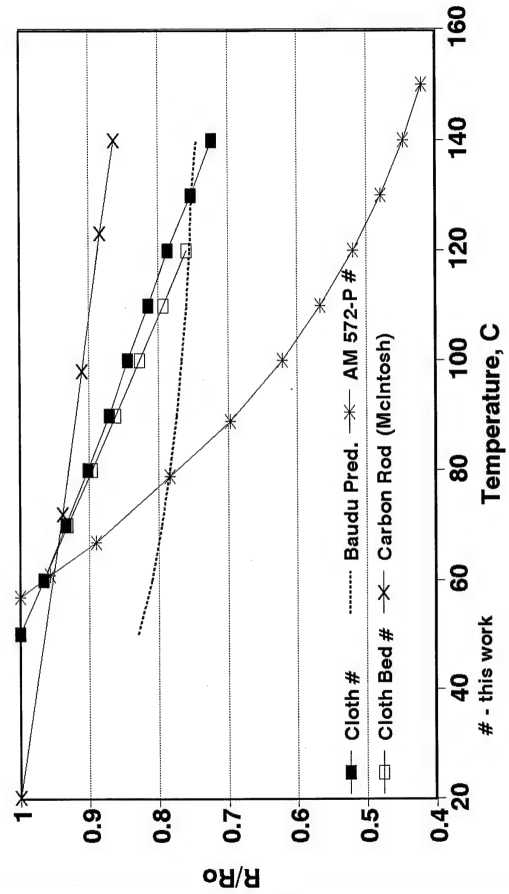


Figure 4-3. Resistance vs Temperature for Carbon Materials

internal pressure, decreasing the electrode/bed contact resistance, artificially lowering the resistance for a given increase in temperature. Using cloth sections stacked up in glass tube between stainless steel screens (the flow through cell for the dynamic isotherms) there was a slight trend in this direction at higher temperatures as shown in Figure 4-3, however, this is not enough to account for the trend predicted.

Fung et al (1991) report the resistance change with temperature over phenolic-based carbon fibers. Over a temperature range of 30 to 300 °K they found the conductivity varied according to Mott's law (Mott, 1987):

$$\sigma(T) = \sigma_0 \exp[-(To/T)^{1/p}]$$

where  $\sigma$  is the conductivity, To is a fitting parameter, and p=2 in their work. At temperatures above ~100 °K, there appears to be a direct linear relationship between the conductivity and the temperature, evidently in the flat tailing portion of the exponential equation. They were unable to definitively say which conduction mechanism governs the electrical transport at room temperature. Later on the same group (Fung et al, 1994) report that the fibers behave like granular metals and a hopping conduction mechanism is dominant in these porous materials.

The overall resistance of the adsorbent bed is very sensitive to pressure and electronic actions. Firm compression of the cloth against the electrode or an electrical burst across the bed (approximately 0.1 s at 42 VDC) dramatically reduces the resistance of the system. All cloth samples were treated with electrical bursts to reduce the overall resistance of the materials prior to use.

The pressure on the carbon cloth cell could be causing three things: 1) improving the fiber:fiber contact, 2) improving the electrode:bed contact, and/or 3) causing the graphitic planes to be making better contact within the cloth. Karanja et al (1994) investigated the pressure dependence of conductivity on pure and doped samples of a polyproplyene polymer film. The material possessed amorphous and crystallite

regions like the carbon fibers. They found that above 2 MPa (20 atm) the conductivities of the materials changed. The pressures here are 2-3 atm (in the compression tests), so the contacts internal to the fiber were neglected.

To investigate the importance of the fiber:fiber contact two experiments were performed. The first one utilized a woven strand (approximately 7 cm long) that was removed from the woven cloth and placed between two glass plates with the strand extending out for direct contact with alligator leads to the volt-ohm meter. Only a slight (0.5 - 1 percent) drop in resistance was noted during compression (ca, 15 psi), however, upon release the conductivity was drastically reduced and in some cases the strand was powderized.

The second experiment to discern the importance of the fiber contacts utilized three strips of the carbon cloth (2 cm by 7 cm). They were stacked up in a star pattern and the central overlapping area of approximately 4 cm² was placed between two glass sheets. Electrical leads were connected to one end of the strip on the bottom and to one end of the strip on the top layer. Upon compression (ca, 30 psi) the resistance dropped by approximately 10 percent and returned to normal when the pressure was released. The experiment was repeated with the leads on opposite ends of the same strip and saw the same result. Thus the fiber:fiber contact is important, and it may not be possible to determine if the electrode:bed or the fiber:fiber contact plays the dominant role. An operating reactor design may utilize large stacks of cloth sections if the majority of the resistance is in the electrode:bed contacts.

The single woven strand, out of its weave, has a resistance on the order of 1000 ohm/cm. A single woven strand, still in its weave (except for the small amount for electrical contact), is approximately 100 ohm/cm. Apparently the direction for the electron flow is not necessarily along the fiber length, but proceeds rather erratically across the material. That is, the preferred electron path, based on the reduced resistance noted, is from one woven fiber to another and not along a single fiber strand. This is

reinforced by the observation in the lab that the resistance from a single point on a carbon woven section is essentially the same  $(\pm 2\%)$  at points in radial directions, but equal in distances, from the starting point. Thus there are no favored paths (such as along the fibers) in the material.

Electrical bursts along these paths improve the conductivity of the carbon cloth samples. At the points of electrical junction from one fiber direction to another, there may be small resistances that are removed by the sudden current flow from the short electrical bursts. Thus, the bursts would improve the conductivity not only between the electrode and the bed, but also within the bed as well.

## 4.3.2 Resistance Change upon Adsorption

As the organics fill in the zones between the crystallites, it is thought they interfere with the electron flow. Experiments confirmed this and showed a concentration dependence as well as a compound dependence in the response of the resistance change. Flow through cells allowed for concentrations to be shifted from low to high, while static isotherms allowed for direct comparison of the different compounds and their effects on the resistance change.

Figure 4-4 shows the results of the dynamic adsorption isotherms using dry streams of trichloroethylene (TCE) performed in the flow through cell. The cloth bed was sequentially loaded with 15, 500, and 1500 ppmv TCE in dry air. The bed was allowed to come to equilibrium before switching to the next higher feed concentration.

There appears to be a small increase in resistance when the cloth is exposed to the 15 ppmv concentration. However, at 500 ppmv the resistance decreases as the surface concentration of TCE increases. Since the 500 ppmv feed is in nitrogen and not air as the other two TCE concentrations, the drop in resistance may be due to the loss of oxygen from the surface. However, in separate experiments with nitrogen and oxygen, the cloth showed no difference in conductivity when switching back and forth between nitrogen and

air carrier gases. Thus the loss of resistance is not due to the change in the carrier gas (oxygen or nitrogen).

At 1500 ppmv there is a marked increase in resistance of the bed. Due to the low adsorbent mass in the bed the breakthrough curve and the resistance-change curve appear to overlap one another. However, in a separate experiment, shown in Figure 4-5, it is shown that the resistance change is at the beginning of the loading and not during breakthrough. Figure 4-5 also demonstrates that the resistance change is not due to the electrode:bed interface since there is not a second sharp increase in resistance at the time of breakthrough.

The downward trend at the early times in Figure 4-5 may be the result of the same phenomenon causing the other downward trends noted above. These trends may be from two sources. The first is from the temperature increase due to the heat of adsorption released when the TCE adsorbs to the surface. At low loadings there is not enough adsorbed to alter the temperature and thus the resistance. At 500 ppmv there may be a sufficient amount adsorbed to increase the temperature (which in turn decreases the resistance). While this may occur initially, the continuing reduced resistance over 20 hours as shown in Figure 4-4 at 500 ppmv indicates that it is not a temperature affect. If it were, then the resistance would gradually return as the bed resumed its normal operating temperature once adsorption ceased.

The second source for the downward trend is thought to be the additive effects to the adsorbent from the pi electrons in the double bond of TCE. If TCE is chemisorbed to the adsorbent then an exchange of electrons occurs and the shift in those extra electrons would alter the charge carrier concentration of the adsorbent material. However, as more and more TCE is adsorbed, a barrier to electron flow may have been set up in the adsorbent, causing the resistance to rise. In Figure 4-5, as the concentration front moves through the bed the leading edge initially lowers the resistance of that section. As the front continues the low resistance zone moves down (and eventually out of) the bed. Behind the bed the highly resistive zone develops. At early times in the loading cycle the

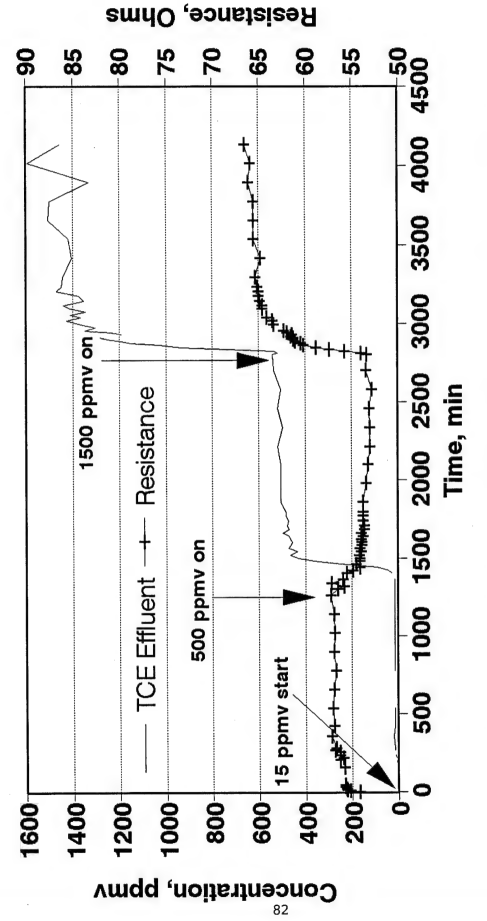


Figure 4-4. Dynamic Isotherm of TCE with Dry Feed (15, 500, and 1500 ppmv) and Resistance Changes on Activated Carbon Cloth (Kuractive)

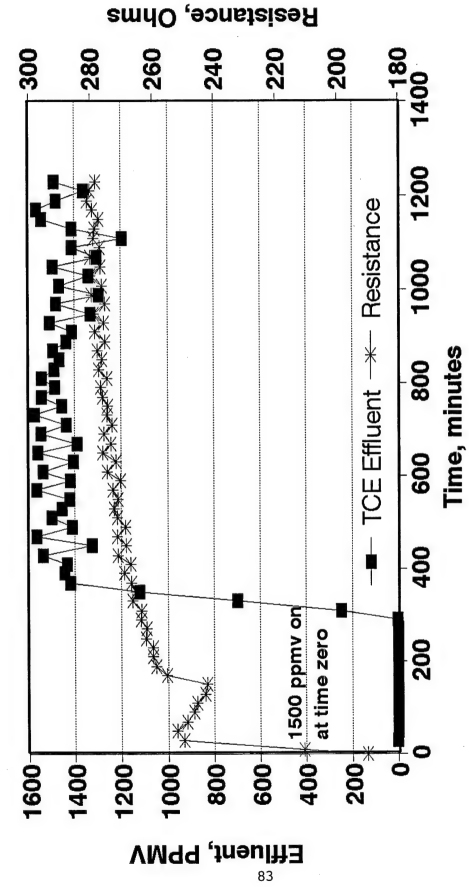


Figure 4-5. 1500 ppmv TCE Humid Feed Loading in Flow Cell over Carbon Cloth

low resistance zone more than balances out the high resistance zone and the overall resistance decreases. As the front moves through the bed the low resistance portion becomes smaller and the high resistance zone dominates for the remainder of the cycle. Further experiments with other double bonded species were conducted to verify the pibond-additive effect in static (no-flow) experiments.

A series of static isothermal experiments were conducted using seven chemicals found in industrial operations or remediation efforts that spanned the range of adsorption strengths, from weak to strong. The compounds are: methanol (MeOH), dichloromethane (DCM), carbon tetrachloride (CCl<sub>4</sub>), tetrachloroethylene (PCE), TCE, water, and toluene. Each experiment used a fresh section of carbon cloth and thus each had a slightly different initial resistance. The changes in resistance were normalized to the initial resistance of cloth sample to allow a comparison between the different compounds. Figures 4-6 and 4-7 show the compounds and their impact on the resistance of the carbon cloth. Since these compounds were exposed to the chemicals at their vapor pressure it is assumed that pore filling, and not just monolayer adsorption, is occurring.

The polarizability of a compound is an indication of how strongly an organic compound will be held to the adsorbent. In addition, the polarizability of the compounds used in this study (listed in Table 4-2) also impacts the timing as to when the resistance change occurs in the adsorbent. Figure 4-6 shows that the weakly adsorbing compounds with the lowest polarizabilities (methanol and dichloromethane) affect the resistance the fastest. Next comes TCE and carbon tetrachloride, which are intermediately adsorbing compounds. The last to show the upward trend in resistance change are the strongly adsorbing PCE and toluene with the largest polarizabilities. The PCE and toluene lines cross making it difficult to determine which comes out later, but both are later than the other compounds.

TCE, PCE and toluene all have double bonds with pi electrons and should behave the same in the adsorbent. However, the initial dramatic downward shift in resistance by TCE is not as strongly demonstrated in the other two compounds. It is

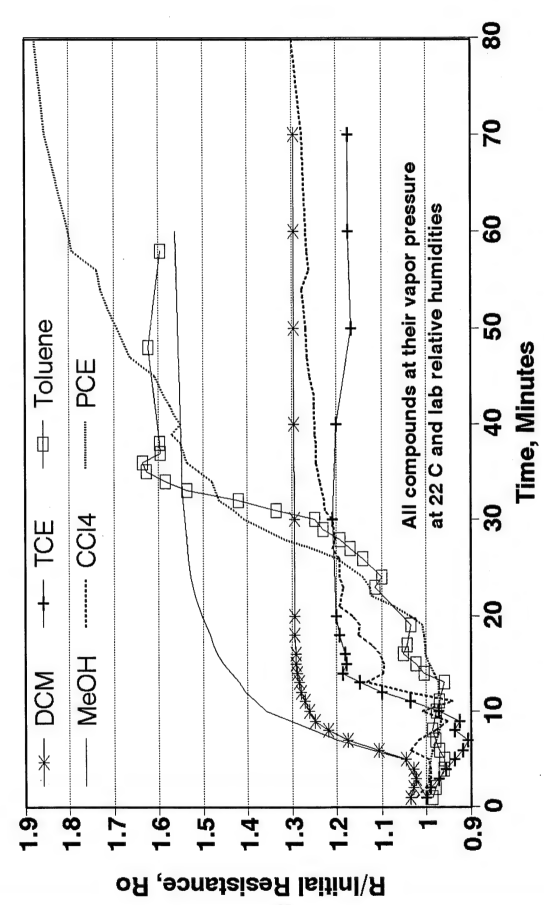


Figure 4-6. Resistance Change Upon Adsorption of Six Compounds on Activated Carbon Cloth (Kuractive)

TABLE 4-2. CHEMICAL CHARACTERISTICS OF SIX COMPOUNDS

COMPOUND	POLARIZ- ABILITY <sup>a</sup> cc-1	MOLAR VOLUME <sup>b</sup> L/mol	VAPOR PRESSURE <sup>c</sup> kPa	O° g/cc	€ <sup>d</sup>
METHANOL	13.3	.042	14.4	1.54	33.6
DICHLORO- METHANE	17.8	.07	51.8	1.70	9.08
TRICHLORO- ETHYLENE	28.7	.102	8.53	1.53	3.4
CARBON TETRACHLORIDE	31.4	.105	13.3	1.58	2.24
TETRACHLORO- ETHYLENE	35.3	.119	2.10	2.01	
TOLUENE	34.6	.119	3.23	.756	2.38

a- calculated from molar refraction equation,  $(n^2 - 1)/(n^2 + 2) * Molar volume$ , "n" is refractive index.

thought that because PCE and toluene are so strongly held by the adsorbent on the way to the micropores, they would take longer to reach the sites altering the resistance of the adsorbent and the influence of their pi electrons may be masked by the blocking mechanism at high concentrations.

In adsorption the adsorbates move by pore and surface diffusion. The driving force for the movement is the concentration of the adsorbate, while the restricting factor is usually the strength of adsorption. If the compound is strongly held then the concentration front moves slowly towards the inner reaches of the adsorbent. However if the compound is weakly held then the concentration front moves as fast as the driving force allows.

Thus, the concentration "front" that moves along the surface toward the high energy sites would be sharper when very strongly or very weakly adsorbing molecules are

b- molar volume from StEPP (Software Estimation of Physical Properties) @ normal boiling point

c-density from StEPP @ 22 °C

d-dielectric constants from CRC Handbook of Chemistry and Physics, 1980

used. This indicates that the PCE and toluene reach the micropores later and in much greater initial concentration than the TCE. With these compounds there is no time for the pi electrons to add to the adsorbate carrier concentration as with TCE. Thus, the downward trend in the resistance is not seen since the pi electron contribution to the matrix during the loading cycle is masked with strongly adsorbing compounds.

The height of the plateaus in Figure 4-6 are perplexing. Since the pores contain condensed organics, one would expect the resistance change would be due to the dielectric constant of the organic. However, the dielectric constants listed in Table 4-2 do not correlate with the height of the curves. Neither does the height relate to the concentrations (vapor pressures), also listed in Table 4-2.

Figure 4-7 shows the resistance change upon adsorption for water, TCE and a water/TCE mixture under static conditions. The mixture contained sufficient contaminant to saturate the cloth and keep both compounds at their vapor pressures, yet the water was not covering the TCE in the bottom of the isotherm bottle. The initial response of the mixture follows the TCE line, however the water does impact the system causing a delay in the upward swing of the resistance curve. The water may have condensed in a sufficient number of pores to delay the transfer of TCE to the high energy sites. However, the resistance change appears to be additive. That is, the initial decrease by the TCE appears to have been enhanced by the initial decrease of the water creating a steeper initial slope in the mixture.

### 4.3.3 Applications

Some useful applications of this finding would be in automatic controls for gas phase regeneration and in capacity predictions. For example, if a large adsorbent bed contained a small cloth resistor near the effluent, one could simply monitor the resistance to determine when the bed was exhausted and required regeneration. While it may be possible to also monitor the extent of the regeneration as well, care must be taken that the

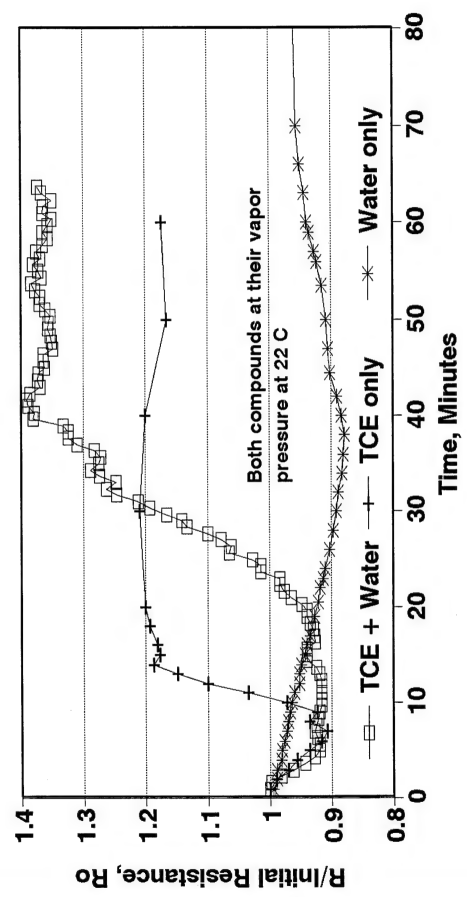


Figure 4-7. Resistance Changes in Mixtures of Trichloroethylene and Water over Carbon Cloth (Kuractive)

resistance change due to the high thermal inputs needed to desorb the organics do not overshadow the resistance change due to desorption.

The other possible application is in determining how a new compound would behave in an existing stream. A static isotherm could be done for the compounds of interest in the existing stream. An additional isotherm could be done with the new compound. If the compound altered the resistance at a later time than the existing stream, then the compound would be held stronger than those compounds. That is, one could predict how strongly or weakly the material would be adsorbed to the adsorbent in relation to the other compounds by knowing the timing of the resistance change in the static isotherms.

#### 4.4 CONCLUSIONS

Neutral organic compounds, in sufficient concentration, can alter the resistance of an activated carbon cloth upon adsorption. The resistance change profile is different for different compounds and for different concentrations of the same compound. At low TCE concentrations (low surface coverage) there is little, if any, resistance change in the design used in this study. Increasing the TCE concentration causes the resistance to drop slightly, while high TCE concentrations increase the resistance significantly. Compounds with large polarizabilities take the longest to cause the upward swing of the resistance curve. These strongly adsorbing compounds are delayed in reaching the areas affecting the resistance change resulting in the delay in the curve. In addition, the conductivity of the adsorbent has a direct linear relationship to temperature. These facts support the concept of the carbon fiber containing crystallites of graphitic-like domains. Electrical conductivity is accomplished by electron hopping from one crystallite to another. Zones between crystallites are the micropores and the adsorption sites are probably at the edge of the basal planes of the crystallites. Thus, adsorption increases the barrier to electron flow and the resistance increases.

Simple tools based on this phenomenon can be employed easily and can reduce monitoring costs, achieve optimum utilization of air phase systems by only regenerating the adsorbent bed when exhausted, and aid engineers in estimating the effects of compounds on the capacity of systems already in use. To expand the usefulness of this work, other compounds and adsorbents should be investigated to further delineate the interactions of the adsorbates and the adsorbent upon adsorption, including multicomponent efforts. In addition other designs need to be evaluated to improve the measurements, especially to define the minimum concentration detection limit for this new process.

#### 4.5 ACKNOWLEDGEMENTS

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# Chapter 5

# Overall Conclusions and Recommendations

# 5.1 OVERALL CONCLUSIONS

Catalytic oxidation over a thermally modified Ambersorb 572 can be accomplished at temperatures as low as 110 °C, although many by-products are formed at that temperature. These by-products include chloroform, carbon tetrachloride, tetrachloroethylene, hexachloroethylene, and, under extremely dry conditions, phosgene. If the temperature is maintained at or above 180 °C then complete mineralization of a 1500 ppmv trichloroethylene (TCE) air stream is possible using plain Ambersorb 572. The addition of transition metal oxides do not enhance the catalytic activity of the TCE oxidation reaction under the conditions used in this research (temperatures below 250 °C). Humidity added to the feed stream deactivates all the catalysts except for the plain Ambersorb 572.

Dynamic electric fields applied to the adsorbents used in this work cause the material to be rapidly heated. Depending on the magnitude of the potential, the heat can 1) alter the local humidity of an adsorbent allowing for increased adsorption capacity of organics; 2) thermally drive off the organics from the surface of the adsorbents during regeneration; and 3) provide the activation energy to catalytically oxidize TCE over an Ambersorb-572 based catalyst.

The electron flow path in the carbon cloth fibers is proposed to travel parallel to the basal planes inside crystallites in the material. Electron hopping, from one crystallite to another provides the overall conductivity. The fringe area of the crystallites are suspected to be the micropores where adsorption occurs. Thus, adsorption at those sites alters the overall resistance of the adsorbent. All the adsorbates used in this work (carbon tetrachloride (CCl<sub>4</sub>), tetrachloroethylene (PCE), TCE, methanol, water, dichloromethane,

and toluene) eventually increased the resistance of the adsorbent. The time for the rapid upswing in resistance change is directly related to the polarizability of the compound. In addition, the concentration of the adsorbate also plays a key role. At low concentrations there may be no effect. Increasing the concentration of TCE caused the resistance to drop, until a higher concentration was used, causing a rapid rise in the resistance of the carbon cloth. Interference, positively and negatively, in the conductivity of the material occurred.

#### 5.2 OVERALL RECOMMENDATIONS

The catalytic oxidation work with TCE should be expanded to other compounds. Since the chloride radical (Cl•) may play a role in the oxidation, non-chlorinated organics should be investigated. In addition, mixture effects should also be evaluated. Long-term tests should be done immediately to determine how long the catalyst material would function on a TCE stream to allow for a life-time cost analysis to be performed.

Initial cost estimates for the using electrical energy to alter the local relative humidity are not favorable and heating of the air should be used. Evaluations should be made to determine if on-site regeneration could be done by exposing the adsorbent to a clean humidified stream at ambient temperatures. Upon restart there would be a small residual until the bed reached the optimum temperature. This time could be reduced by using the Joule heating to quickly heat the bed, then switching it off for the remainder of the cycle. Evaluations with Joule heating should be done to optimize the adsorption capacity for organics by looking at the competing terms: reduced organic capacity at elevated temperatures and reduced relative humidity at increased temperatures.

The resistance change upon adsorption phenomenon should be evaluated for other adsorbents. Once an optimum one is found, then it should be applied in the field monitoring air. For example a unit could monitor the effluent from an air stripping operation to determine when the gas phase adsorption unit needed regeneration. Another possible application is between the walls of double-walled gas tanks to monitor for leaks.

In addition, single solute and mixtures should be used on different adsorbents and compared to observed time to breakthrough from long-term isotherm studies. These simple tests could then be used to strengthen the database of chemical adsorptabilities onto different adsorbents. These experiments could be done economically and without lengthy isotherm tests.

# APPENDIX A

Raw Data from Adsorption/Desorption Data from PMI Inc., (Ithaca, NY) for Ambersorb 572 & MTU-CAT

#### BET SURFACE AREA ANALYSIS

### 12-16-1994

POROUS MATERIALS, INC. ANALYTICAL SERVICES DIVISION CORNELL INDUSTRY RESEARCH PARK, BLDG. 4 ITHACA, NY 14850 USA PHONE (607) 257-4267 OR 257-5544

NOTE: RESULTS CALCULATED USING WEIGHT AFTER OUTGASSING. TO GET RESULTS USING ORIGINAL WEIGHT, MULTIPLY THESE VALUES BY WT AFTER OUTGAS/WT BEFORE OUTGAS

MTU

Ed. Marchand

SAMPLE ID: Ambersor 572

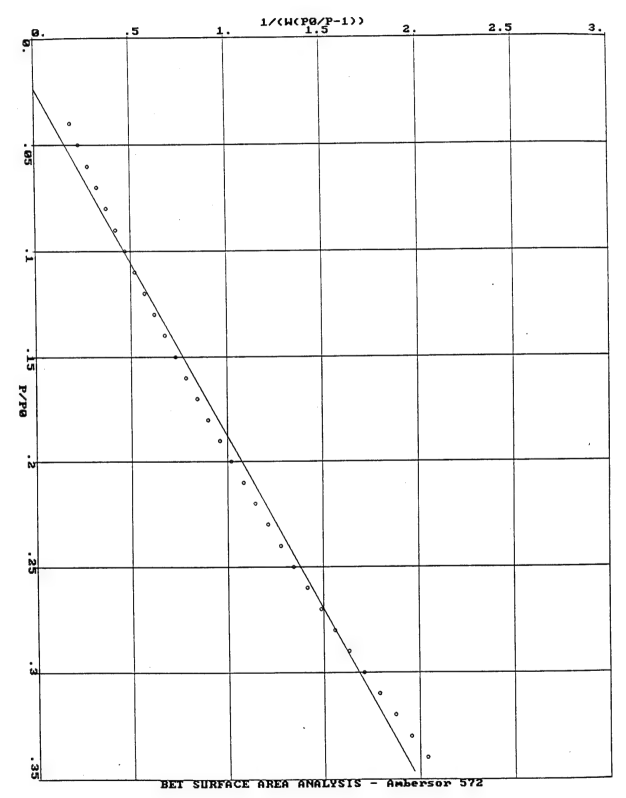
## SPECIFIC SURFACE AREA= 830.9971 M2/GM

SAMPLE WEIGHT= .7196 GM BEFORE OUTGASSING .703 GM AFTER OUTGASSING Sample Density= 1.5
BET C VALUE= -40.612
SLOPE= 6.107 GM-1
Y INTERCEPT= -.147 GM-1
CORRELATION COEFFICIENT= .995
ADSORBATE= NITROGEN
VA VOLUME= 44.80148 CC
Vit VOLUME= 11.7589 CC
Vln VOLUME= 13.782 CC
INSTRUMENT TEMPERATURE= 305.36 K
ROOM TEMPERATURE= 296.7 K
W = Wt. of adsorbed gas per gram of sample

SAMPLE OUTGASSED AT 20 DEGREES CELSIUS

	P	PO	-1
DATA	- (W	I(1)	))
POINT	PO `	`P	•
9	0.04000000	0.186	00000
10	0.05000000	0.23	100000
11	0.06000000		700000
12	0.07000000		500000
13	0.0800000		300000
14	0.09000001		100000
15	0.10000000		900000
16	0.11000000		900000
17	0.12000000		00000
18	0.13000000		200000
19	0.14000000		00000
20	0.15000000		00000
21	0.16000000		00000
22	0.17000000		100000
23	0.18000000		00000
24	0.19000000		300000
		44330	,,,,,,,,

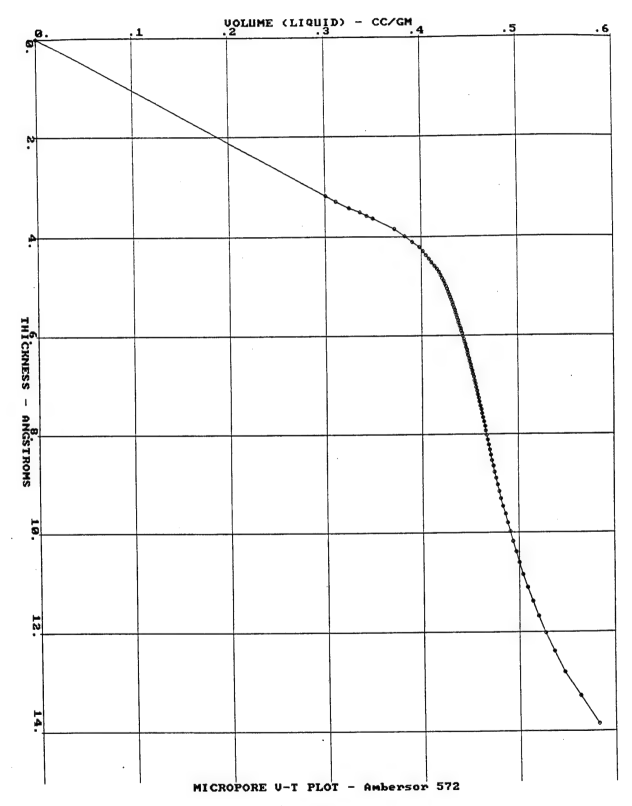
25	0.2000000	1.019
26	0.21000000	1.081
27	0.2200000	1.144
28	0.23000000	1.209
29	0.24000000	1.275
30	0.2500000	1.343
31	0.26000000	1.413
32	0.2700000	1.485
33	0.28000000	1.558
34	0.2900000	1.634
35	0.3000000	1.711
36	0.31000000	1.79
37	0.32000000	1.871
38	0.33000000	1.955
39	0.34000000	2.041



## MICRO PORE CALCULATIONS

Thickness Range Angstroms	Volume (Liquid) cc/g	Hydraulic Radius Angstroms
3.180 - 3.292 3.292 - 3.425 3.425 - 3.513 3.513 - 3.581 3.581 - 3.638 3.638 - 3.842 3.842 - 3.984 3.984 - 4.100 4.100 - 4.199 4.199 - 4.288 4.288 - 4.369 4.369 - 4.445 4.445 - 4.516 4.516 - 4.584 4.584 - 4.649 4.649 - 4.712 4.773 - 4.832 4.832 - 4.890 4.890 - 4.947 4.947 - 5.002 5.002 - 5.057 5.057 - 5.112 5.165 - 5.219 5.219 - 5.272 5.272 - 5.324 5.324 - 5.377 5.377 - 5.429 5.429 - 5.481 5.481 - 5.533 5.533 - 5.585 5.585 - 5.638 5.638 - 5.690 5.690 - 5.743 5.743 - 5.902 5.902 - 5.956 6.010 - 6.065 6.065 - 6.120 6.176 - 6.232 6.232 - 6.289 6.347 - 6.465 6.465 - 6.525 6.525 - 6.586		
6.586 - 6.648 6.648 - 6.711 6.711 - 6.775 6.775 - 6.840	0.451 0.452 0.453 0.453	6.617 6.679 6.743 6.807

6.840	_	6.906	0.454	6.873
6.906	_	6.974	0.455	6.940
6.974	_	7.043	0.456	7.009
7.043	_	7.114	0.457	7.078
7.114	_	7.186	0.457	7.150
7.186	_	7.259	0.458	7.222
7.259	_	7.335	0.459	7.297
7.335	-	7.412	0.460	7.373
7.412	_	7.491	0.461	7.452
7.491	_	7.572	0.462	7.532
7.572	-	7.656	0.462	7.614
7.656	_	7.742	0.463	7.699
7.742	_	7.830	0.464	7.786
7.830	_	7.921	0.465	7.876
7.921	_	8.015	0.466	7.968
8.015	-	8.112	0.467	8.064
8.112	_	8.212	0.468	8.162
8.212	-	8.316	0.469	8.264
8.316	_	8.424	0.471	8.370
8.424	_	8.536	0.472	8.480
8.536	-	8.652	0.473	8.594
8.652	_	8.773	0.474	8.712
8.773	-	8.899	0.475	8.836
8.899	-	9.031	0.477	8.965
9.031	-	9.170	0.478	9.101
9.170	-	9.315	0.480	9.242
9.315	-	9.468	0.482	9.391
9.468	-	9.629	0.484	9.548
9.629	-	9.799	0.486	9.714
9.799	-	9.980	0.488	9.890
9.980	-	10.172	0.491	10.076
10.172	-	10.378	0.494	10.275
10.378	-	10.598	0.497	10.488
10.598	-	10.836	0.500	10.717
10.836	-	11.092	0.504	10.964
11.092	-	11.372	0.508	11.232
11.372	-	11.678	0.514	11.525
11.678	-	12.017	0.520	11.847
12.017	-	12.393	0.527	12.205
12.393	-	12.816	0.537	12.605
12.816	-	13.299	0.547	13.058
13.299	-	13.856	0.563	13.577



### T-PLOT ANALYSIS:

T-PLOT INTERCEPT: T-PLOT SLOPE: 0.35334 0.013941

MICROPORE VOLUME:

0.35334 CC/GM

FILM THICKNESS USED BETWEEN 8.000 and 12.000 ANGSTROMS

SURFACE AREA = 139.4092 SQ. M/GM (MESO AND MACROPORES ONLY)

MICROPORE SURFACE AREA = 691.5879 SQ. M/GM

## GAS ADSORPTION/DESORPTION RESULTS

ADSORBATE: NITROGEN

## ISOTHERM AND PORE VOLUME DISTRIBUTION

## ADSORPTION

	PORE	VOLUME	LIQ. VOL	AVE.	CUMUL.	Δv	∆s	SURFACE	CUMUL.
P/P0	DIAMETER	ADSORBED	ADSORBED I					AREA S	. AREA
-,	Å	CC/GM STP	CC/GM	Å	CC/GM	ďΔ	Δp	M2/GM	M2/GM
0.994	3246.4	586.3235	0.90294	3246.43	-0-	1.79E-06	2.75E-05	-0-	-0-
0.990	1957.7	584.8974	0.90074	2602.07	0.00230	2.90E-06	6.55E-05	0.03541	0.03541
0.985	1313.5	583.5333	0.89864	1635.60	0.00453	1.56E-05	5.30E-04	0.05451	0.08992
0.980	990.44	579.3146	0.89214	1151.97	0.01157	7.70E-05	0.00338	0.24430	0.33422
0.975	796.1	566.6119	0.87258	893.270	0.03309	2.08E-04	0.01095	0.96380	1.29802
0.970	666.21	545.8799	0.84066	731.155	0.06864	3.78E-04	0.02325	1.94490	3.24292
0.965	573.22	521.5602	0.80320	619.717	0.11075	5.40E-04	0.03820	2.71803	5.96095
0.960	503.33	497.3028	0.76585	538.275	0.15309	6.85E-04	0.05500	3.14639	9.1073
0.955	448.85	473.9495	0.72988	476.088	0.19416	7.90E-04	0.07050	3.45025	12.5576
0.950	405.18	453.6172	0.69857	427.013	0.23009	8.00E-04	0.07900	3.36560	15.9232
0.945	369.37	437.7131	0.67408	387.274	0.25821	7.95E-04	0.08650	2.90468	18.8279
0.940	339.48	424.0804	0.65308	354.426	0.28238	8.80E-04	0.10450	2.72815	21.5560
0.930	292.37	397.8874	0.61275	315.927	0.32948	0.00100	0.13700	5.96352	27.5195
0.920	256.92	378.2267	0.58247	274.646	0.36489	8.75E-04	0.13600	5.15686	32.6764
0.910	229.24	365.8985	0.56348	243.079	0.38664	8.15E-04	0.14350	3.57956	36.2559
0.900	207.02	355.3108	0.54718	218.133	0.40543	7.30E-04	0.14100	3.44483	39.7008
0.890	188.79	348.4164	0.53656	197.905	0.41717	6.65E-04	0.14150	2.37196	42.0727
0.880	173.54	342.307	0.52715	181.161	0.42762	6.35E-04	0.14600	2.30761	44.3804
0.870	160.59	337.6506	0.51998	167.063	0.43532	5.65E-04	0.14100	1.84345	46.2238
0.860	149.46	333.8754	0.51417	155.025		5.90E-04	0.15850	1.56571	47.7895
0.850	139.78	330.177	0.50847	144.619	0.44749	5.85E-04	0.16750	1.68994	49.4795
0.840	131.28	327.2249	0.50393	135.532		5.05E-04	0.15400	1.37580	50.8553
0.830	123.76	324.8524	0.50027	127.524		4.93E-04	0.15950	1.10893	51.9642
0.820	117.06	322.58	0.49677	120.412	0.45914	4.87E-04	0.16600	1.14699	53.1112
0.810	111.04	320.6564	0.49381	114.050	0.46193	4.74E-04	0.17050		54.0875
0.800	105.61	318.8483	0.49103	108.323	0.46456	4.87E-04	0.18400	0.97198	55.059 <b>5</b>
0.790	100.67	317.1794	0.48846	103.141		4.77E-04	0.18950	0.93491	55.9944
0.780	96.177	315.6809	0.48615	98.4258		4.96E-04	0.20650	0.85384	56.8483
0.770	92.057	314.1973		94.1170		5.15E-04	0.22450	0.91555	57.7638
0.760	88.269	312.8355	0.48177	90.1629		4.89E-04		0.85978	58.6236
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81.534 310.5025 0.47817 83.1529 0.47629 4.43E-04 0.21700 0.71524 60.0962
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0.730
                            0.47515 77.1244 0.47873 3.60E-04 0.18900 0.59949 61.3384
0.720
         75.723 308.54
        73.10501 307.7486 0.47393 74.4142 0.47954 3.19E-04 0.17450 0.43631 61.7748
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                  306.2431 0.47161 69.5029 0.48112 3.46E-04 0.20250 0.44389 62.6646
0.690
         68.352
                  305.5225 0.47050 67.2697 0.48189 3.78E-04 0.22850 0.45852 63.1232
0.680
         66.187
                           0.46940 65.1668 0.48270 4.15E-04 0.25850 0.49982 63.6230
0.670
         64.14601 304.8023
                  0.660
         62.219
0.650
         60.395
0.640
         58.666
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0.630
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                           0.46440 56.2433 0.48635 4.97E-04 0.35900 0.52605 66.0693
0.620
         55.463
                  301.5591
0.610
                300.9316
                           0.46343 54.7192 0.48712 5.35E-04 0.39700 0.56513 66.6344
         53.976
                300.3117 0.46248 53.2668 0.48790 5.10E-04 0.39000 0.58661 67.2210
0.600
         52.558
         51.203 299.7455 0.46161 51.8806 0.48855 4.68E-04 0.36550 0.49850 67.7195
0.590
                299.2052 0.46078 50.5559 0.48914 4.80E-04 0.38450 0.47054 68.1901
0.580
        49.908
        48.668 298.6655 0.45994 49.2884 0.48976 5.20E-04 0.42950 0.50301 68.6931
0.570
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0.560
        47.48
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0.550
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        45.243 297.0625 0.45748 45.7909 0.49172 5.90E-04 0.52500 0.56304 70.3625
0.540
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0.530
                                                                               70.9277
                296.0352 0.45589 43.6812 0.49300 6.70E-04 0.62000 0.59765
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0.470
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                0.410
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0.400
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0.370
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0.360
0.350
        30.116 287.3629 0.44254 30.4217 0.50633 0.00159 2.11000 1.20706 86.5412
                 286.8247 0.44171 29.8181 0.50733 0.00171 2.31000 1.34219 87.8834
0.340
        29.52
0.330
        28.936 286.2839 0.44088 29.2277 0.50834 0.00176 2.43000 1.38952 89.2729

    0.44004
    28.6498
    0.50936
    0.00181
    2.55000
    1.42128
    90.6942

    0.43921
    28.0837
    0.51039
    0.00185
    2.66500
    1.46456
    92.1587

    0.43837
    27.5287
    0.51142
    0.00189
    2.77000
    1.49606
    93.6548

    0.43753
    26.9841
    0.51245
    0.00192
    2.87000
    1.52284
    95.1776

0.320
        28.364 285.743
0.310
         27.804 285.2
                 284.6566
0.300
         27.254
        26.714
0.290
                  284.1132
                 283.5696
                             0.43670 26.4492 0.51347 0.00194 2.96500 1.54909 96.7267
0.280
         26.184
                            0.43586 25.9234 0.51449 0.00197 3.06500 1.56966 98.2964
                 283.0263
0.270
         25.663
        25.149 282.481
                             0.43502 25.4061 0.51551 0.00208 3.30500 1.60455
0.260
                                                                               99.901
                            0.43415 24.8966 0.51660 0.00224 3.63000 1.76136 101.662
0.250
        24.644 281.9178
         24.145 281.3411
0.240
                            0.43327 24.3943 0.51775 0.00230 3.81500 1.88525 103.548
0.230
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         23.652 280.7639
0.220
                             0.43145 23.4087 0.52013 0.00301 5.20000 2.11817 107.566
         23.165 280.1624
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         22.683
                  279.4735
0.200
         22.205
                  278.7612
                            0.42929 22.4443 0.52356 0.00365 6.55000 3.13770 113.621
0.190
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                   278.049
                             0.42820 21.9683 0.52528 0.00358 6.55000 3.12582 116.746
0.180
         21.26
                             0.42710 21.4954 0.52695 0.00349 6.55000 3.10445 119.851
                   277.3366
                             0.42600 21.0250 0.52856 0.00375 7.20000 3.07375 122.925
0.170
         20.79
                   276.6234
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0.160
         20.322
                   275.8499
0.150
                             0.42335 20.0878 0.53331 0.00605 12.1000 5.67748 132.296
         19.854
                  274.9012
```

0.140	273.9191
0.130	272.937
0.120	271.9488
0.110	270.6335
0.100	268.7639
0.0900	266.8271
0.0800	264.8903
0.0700	262.9529
0.0600	261.0071
0.0500	258.3368
0.0400	253.5057
0.0300	248.2402
0.0200	241.3405
0.0100	227.008
0.00800	222.928
0.00600	218.1931
0.00400	211.1391
0.00200	202.0771
0.00101	195 2435

### SUMMARY SHEET

UTM

Ed. Marchand

SAMPLE ID: Ambersor 572

Total Surface Area = 830.9971 sq. m/gm

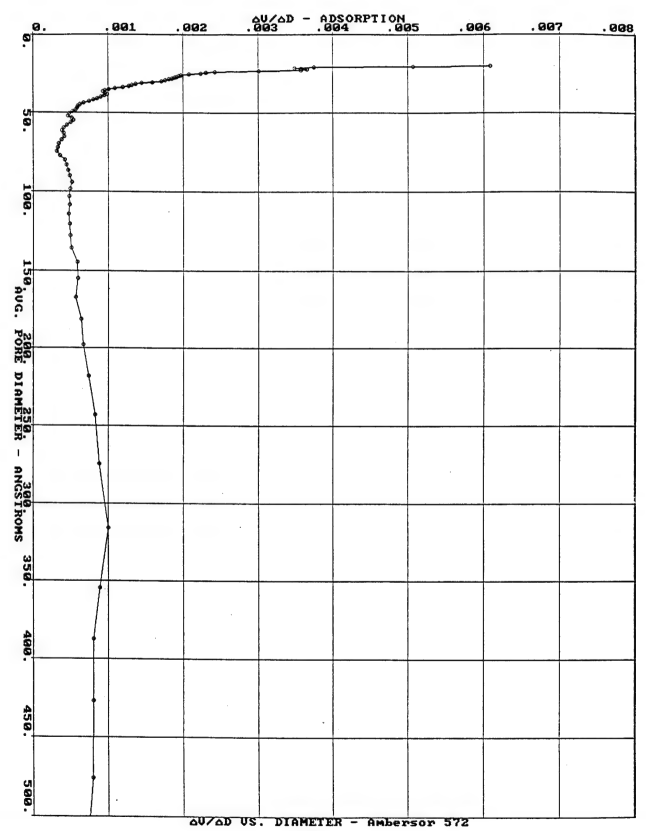
Average Pore Diameter (4V/S) = 43.6559 Angstroms

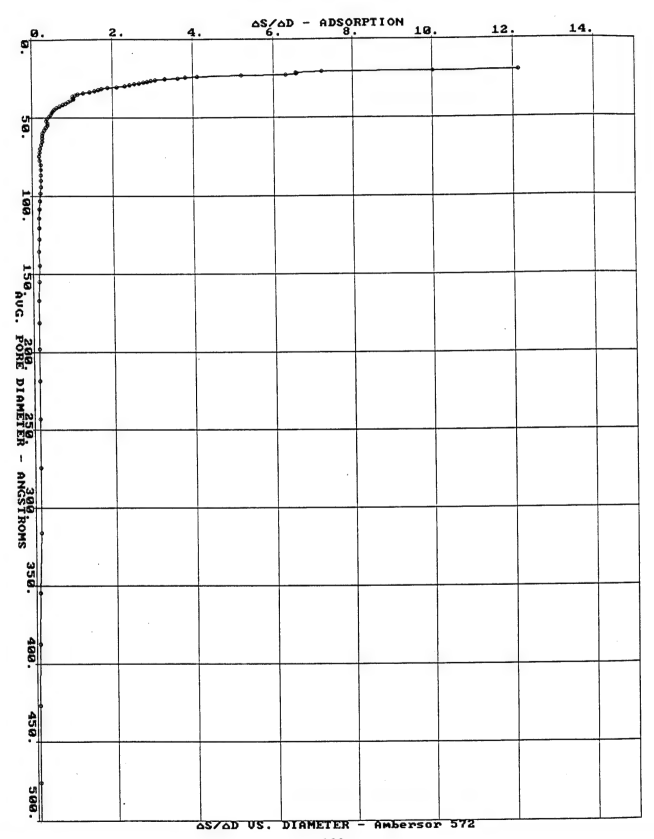
Total Pore Volume = 0.9069 cc/g

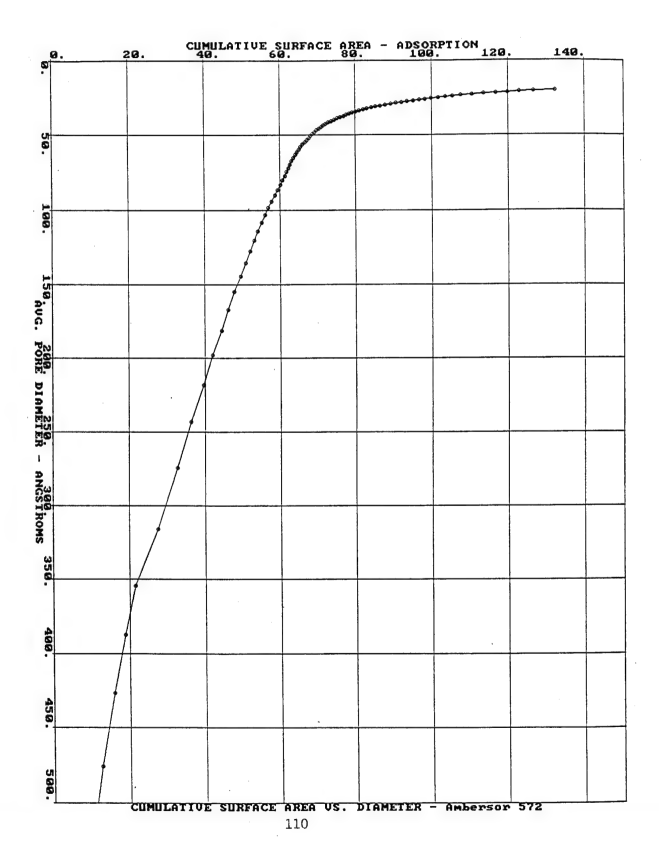
Median Pore Diameter (based on pore volume) = 132.547 Angstroms
Standard Deviation = 72.8709 Angstroms

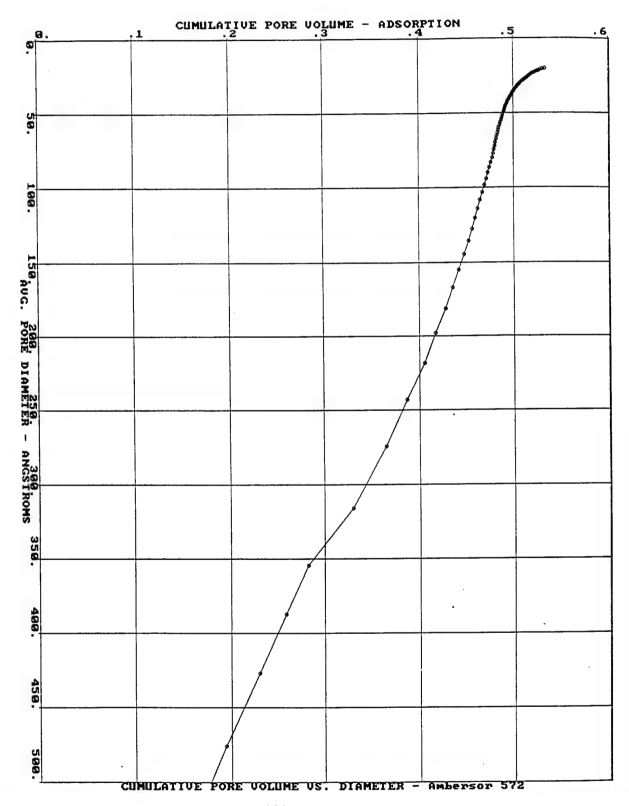
Median Pore Diameter (based on surface area) = 56.03 Angstroms

Standard Deviation = 22.5873 Angstroms









## GAS ADSORPTION/DESORPTION RESULTS

ADSORBATE: NITROGEN

## ISOTHERM AND PORE VOLUME DISTRIBUTION

## DESORPTION

	PORE	VOLUME	LIQ. VOL	AVE.	CUMUL.	Δv	∆s	SURFACE	CUMUL.
P/P0	DIAMETER	ADSORBED	ADSORBED P	OR.DIAM	POR. VOL			AREA S	. AREA
	Ā	CC/GM STP	CC/GM	Ā	CC/GM	Δp	∆¤	M2/GM	M2/GM
0.990	1957.7	585.5026	0.90167	1957.70	-0-	2.12E-06	5.15E-05	-0-	-0-
0.985	1313.5	584.6721	0.90040	1635.60	0.00137	2.38E-06	7.50E-05	0.03339	0.03339
0.980	990.44	584.1822	0.89964	1151.97	0.00218	2.54E-06	1.04E-04	0.02818	0.06156
0.975	796.1	583.8838	0.89918	893.270	0.00267	3.02E-06	1.55E-04	0.02222	0.08378
0.970	666.21	583.626	0.89878	731.155	0.00311	4.14E-06	2.54E-04	0.02368	0.10746
0.965	573.22	583.3671	0.89839	619.717	0.00355	6.00E-06	4.26E-04	0.02840	0.13585
0.960	503.33	583.0847	0.89795	538.275	0.00403	9.15E-06	7.40E-04	0.03617	0.17202
0.955	448.85	582.745	0.89743	476.088	0.00463	2.84E-05	0.00261	0.04992	0.22195
0.950	405.18	581.7145	0.89584	427.013	0.00648	2.16E-04	0.02210	0.17331	0.39525
0.945	369.37	574.6946	0.88503	387.274	0.01930	8.70E-04	0.09700	1.32471	1.71996
0.940	339.48	553.5632	0.85249	354.426	0.05830	0.00177	0.21300	4.40164	6.12160
0.930	292.37	490.2859	0.75504	315.927	0.17653	0.00300	0.41650	14.9687	21.0903
0.920	256.92	426.907	0.65744	274.646	0.29619	0.00281	0.43450	17.4279	38.5182
0.910	229.24	391.8633	0.60347	243.079	0.36186	0.00164	0.28000	10.8070	49.3252
0.900	207.02	378.6347	0.58310	218.133	0.38509	0.00113	0.22000	4.25814	53.5833
0.890	188.79	366.2789	0.56407	197.905	0.40709	0.00134	0.28500	4.44838	58.0317
0.880	173.54	354.1259	0.54535	181.161	0.42917	0.00136	0.31250	4.87527	62.9070
0.870	160.59	344.7998	0.53099	167.063	0.44579	0.00100	0.24700	3.97776	66.8847
0.860	149.46	339.5018	0.52283	155.025	0.45422	6.85E-04	0.18300	2.17529	69.0600
0.850	139.78	335.4441	0.51658	144.619	0.46027	6.50E-04	0.18650	1.67387	70.7339
0.840	131.28	331.6517	0.51074	135.532		6.15E-04	0.18700	1.69005	72.4239
0.830	123.76	328.6333	0.50610	127.524	0.47023	5.80E-04	0.18800	1.32774	73.7517
0.820	117.06	325.8041	0.50174	120.412	0.47424	6.40E-04	0.22000	1.33084	75.0825
0.810	111.04	323.0048	0.49743	114.050		6.35E-04	0.23000	1.44456	76.5271
0.800	105.61	320.6497	0.49380	108.323	0.48161	5.35E-04	0.20350	1.20088	77.7280
0.790	100.67	318.7209	0.49083	103.141	0.48400	4.66E-04	0.18500	0.92927	78.6572
0.780	96.177	317.007	0.48819	98.4258	0.48602	4.09E-04	0.17000	0.81944	79.4767
0.770	92.057	315.5401	0.48593	94.1170		3.92E-04	0.17050	0.65442	80.1311
0.760	88.269	314.1149	0.48374	90.1629	0.48911	3.98E-04	0.18000	0.68703	80.8181
0.750	84.77201	312.8078	0.48172	86.5203	0.49047	4.01E-04	0.18950	0.62757	81.4457
0.740	81.534	311.546	0.47978	83.1529	0.49180	3.83E-04	0.18750	0.64374	82.0894

```
0.47805 80.0297 0.49287 3.43E-04 0.17450 0.53398 82.6234
                   310.4243
0.730
         78.526
                             0.47646 77.1244 0.49380 3.34E-04 0.17600 0.48235 83.1058
         75.723
                   309.3876
0.720
                             0.47494 74.4142 0.49468 2.56E-04 0.13900 0.47209 83.5778
0.710
         73.10501
                   308.3997
                             0.47371 71.8794 0.49512 1.72E-04 0.09700 0.24706 83.8249
0.700
         70.654
                   307.6019
                             0.47256 69.5029 0.49550 1.82E-04 0.10650 0.21574 84.0406
0.690
         68.352
                   306.8543
                             0.47141 67.2697 0.49593 2.22E-04 0.13400 0.25833 84.2990
0.680
         66.187
                   306.1076
                             0.47026 65.1668 0.49643 2.64E-04 0.16450 0.30336
                                                                                 84.6023
0.670
         64.14601 305.3604
                             0.46911 63.1825 0.49698 3.06E-04 0.19700 0.34697
                                                                                 84.9493
0.660
         62.219
                   304.6137
                             0.46796 61.3067 0.49757 3.11E-04 0.20550 0.38947
                                                                                 85.3388
0.650
         60.395
                   303.8682
                             0.46688 59.5301 0.49808 2.60E-04 0.17650 0.34361 85.6824
0.640
         58.666
                   303.1697
                             0.46590 57.8447 0.49846 2.41E-04 0.16950 0.25655
                                                                                 85.9389
         57.024
0.630
                   302.5356
                             0.46494 56.2433 0.49886 2.73E-04 0.19700 0.28435
                                                                                 86.2233
                   301.9068
0.620
         55.463
                                     54.7192 0.49928 2.68E-04 0.19850 0.31411 86.5374
0.610
         53.976
                   301.2821
                             0.46397
                                     53.2668 0.49964 2.43E-04 0.18500 0.26436 86.8017
0.600
         52.558
                   300.6964
                             0.46307
                             0.46220 51.8806 0.49996 2.58E-04 0.20100 0.24925 87.0510
0.590
         51.203
                   300.1326
                             0.46134 50.5559 0.50032 2.96E-04 0.23750 0.28253 87.3335
0.580
         49.908
                   299.5696
                             0.46047 49.2884 0.50071 3.37E-04 0.27700 0.31800 87.6515
0.570
         48.668
                   299.0066
                             0.45960 48.0741 0.50113 3.76E-04 0.31700 0.35375 88.0053
                   298.4435
0.560
         47.48
                             0.45874 46.9094 0.50158 3.73E-04 0.32200 0.38377 88.3891
0.550
         46.339
                   297.8825
                            0.45792 45.7909 0.50197 3.35E-04 0.29600 0.33791
                                                                                 88.7270
0.540
         45.243
                   297.3502
                            0.45714 44.7158 0.50231 3.37E-04 0.30450 0.30041
                                                                                 89.0274
0.530
         44.189
                   296.8421
                             0.45636 43.6812 0.50267 3.75E-04 0.34750 0.32893
                                                                                 89.3563
0.520
         43.174
                   296.3353
                             0.45558 42.6846 0.50305 4.15E-04 0.39350 0.36229 89.7186
                   295.8282
0.510
         42.196
                             0.45480 41.7237 0.50346 4.69E-04 0.45500 0.39312 90.1117
0.500
         41.252
                   295.3216
                             0.45400 40.7962 0.50392 6.40E-04 0.64000 0.44848 90.5602
0.490
         40.341
                   294.8083
                             0.45312 39.9003 0.50461 8.90E-04 0.90500 0.69135 91.2515
                   294.2354
0.480
         39.46
                             0.45218 39.0339 0.50546 0.00102 1.06000 0.87475 92.1263
0.470 .
         38.608
                   293.6225
                             0.45124 38.1954 0.50633 0.00092 0.97000 0.90300 93.0293
0.460
         37.783
                   293.0128
                             0.45040 37.3833 0.50697 6.95E-04 0.75000 0.68548 93.7148
0.450
         36.984
                   292.469
                            0.44965 36.5959 0.50743 6.10E-04 0.67500 0.50510 94.2199
0.440
         36.208
                   291.9785
                             0.44889 35.8319 0.50790 6.45E-04 0.73000 0.52810 94.7480
0.430
         35.456
                   291.4899
                             0.44814 35.0901 0.50839 6.90E-04 0.79500 0.55710 95.3050
0.420
         34.725
                   291.0017
                            0.44739 34.3691 0.50890 7.30E-04 0.86000 0.58850 95.8936
         34.014
                   290.5133
0.410
                             0.44664 33.6679 0.50942 7.80E-04 0.93500 0.61832 96.5119
0.400
         33.322
                   290.025
                             0.44588 32.9853 0.50996 8.45E-04 1.04000 0.66191 97.1738
0.390
         32.648
                   289.5338
                             0.44512 32.3203 0.51055 0.00091 1.13500 0.72161 97.8954
0.380
         31.992
                   289.0366
                             0.44435 31.6720 0.51114 0.00095 1.21500 0.75470 98.6501
0.370
          31.352
                   288.5387
                             0.44358 31.0394 0.51175 0.00099 1.29500 0.78548
                                                                                 99.436
0.360
          30.727
                   288.0405
                            0.44282 30.4217 0.51237 0.00104 1.38500 0.81337 100.249
                   287.5424
0.350
          30.116
                            0.44204 29.8181 0.51301 0.00115 1.56000 0.85802 101.107
                   287.0413
0.340
         29.52
                            0.44125 29.2277 0.51373 0.00127 1.76000 0.98510 102.092
0.330
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                   286.5232
                            0.44044 28.6498 0.51448 0.00133 1.87500 1.04875 103.141
0.320
          28.364
                   285.9991
                             0.43963 28.0837 0.51524 0.00136 1.96000 1.07661 104.217
0.310
          27.804
                   285.4748
                             0.43882 27.5287 0.51599 0.00143 2.10000 1.09975
0.43800 26.9841 0.51680 0.00165 2.47500 1.19070
                                                                                 105.317
0.300
          27.254
                   284.9507
          26.714
                                                                                 106.508
                   284.4166
0.290
                                                       0.00187 2.85000 1.45374 107.962
                              0.43713 26.4492 0.51776
0.280
          26.184
                   283.8481
                              0.43624 25.9234 0.51876 0.00193 3.00000 1.54501 109.507
0.270
          25.663
                   283.2701
                             0.43535 25.4061 0.51975 0.00196 3.12000 1.56322 111.070
0.260
          25.149
                   282.6923
                             0.43445 24.8966 0.52076 0.00210 3.41000 1.61969 112.690
0.250
          24.644
                   282.1093
                             0.43352 24.3943 0.52186 0.00224 3.70500 1.80621 114.496
0.240
          24.145
                   281.5049
                             0.43258 23.8985 0.52298 0.00226 3.82000 1.87106 116.367
0.230
          23.652
                   280.8939
                             0.43164 23.4087 0.52408 0.00224 3.86500 1.87410 118.241
0.220
          23.165
                   280.2829
                            0.43069 22.9242 0.52515 0.00235 4.14000 1.87118 120.112
0.210
          22.683
                   279.6718
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0.200
          22.205
                   279.035
                            0.42860 21.9683 0.52796 0.00352 6.45000 2.95973 125.175
0.190
          21.731
                   278.3096
                              0.42744 21.4954 0.52966 0.00356 6.70000 3.17444 128.350
0.180
                   277.5607
          21.26
                             0.42629 21.0250 0.53131 0.00344 6.60000 3.13156
                                                                                 131.481
0.170
          20.79
                   276.8118
                                                                                 134.558
0.160
          20.322
                   276.0622
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                              0.42388 20.0878 0.53476 0.00400 7.95000 3.73042 138.288
0.150
          19.854
                   275.2496
0.140
                   274.2463
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0.130	273.199
0.120	272.1515
0.110	271.103
0.100	269.8837
0.0900	268.0038
0.0800	265.9261
0.0700	263.8468
0.0600	261.7609
0.0500	258.9777
0.0400	253.9745
0.0300	248.4888
0.0200	241.6162
0.0100	228.3449
0.00800	224.6267
0.00600	220.2782
0.00400	214.0376
0.00200	206.0367
0.00100	199 9052

### SUMMARY SHEET

MTU

Ed. Marchand

SAMPLE ID: Ambersor 572

Total Surface Area = 830.9971 sq. m/gm

Average Pore Diameter (4V/S) = 43.5948 Angstroms

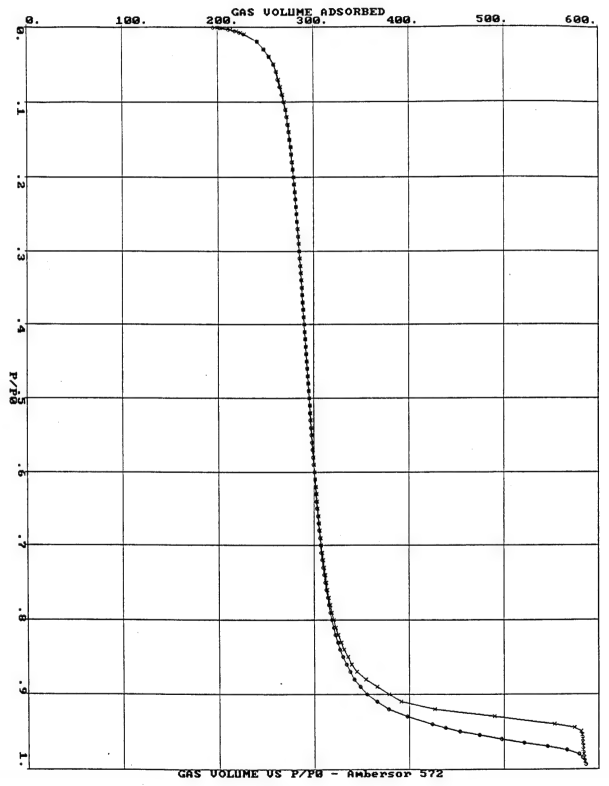
Total Pore Volume = 0.9057 cc/g

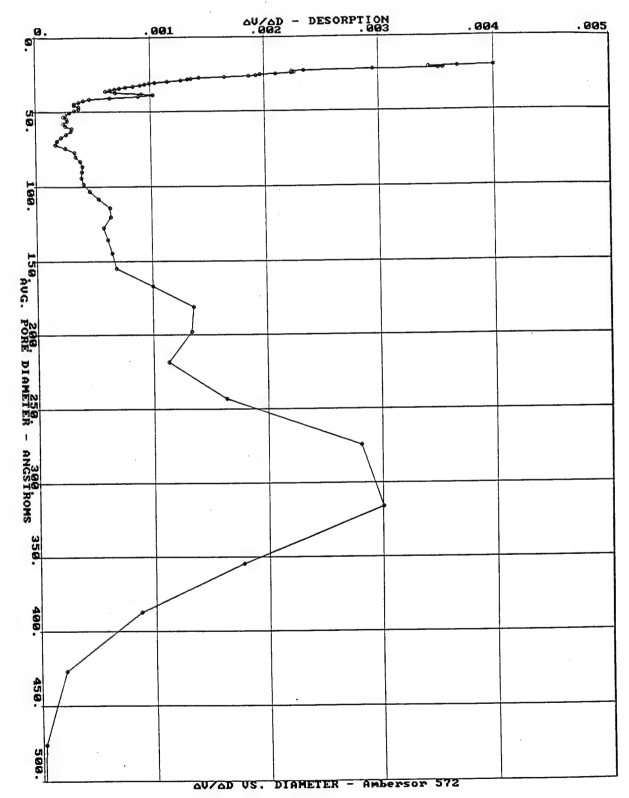
Median Pore Diameter (based on pore volume) = 156.995 Angstroms

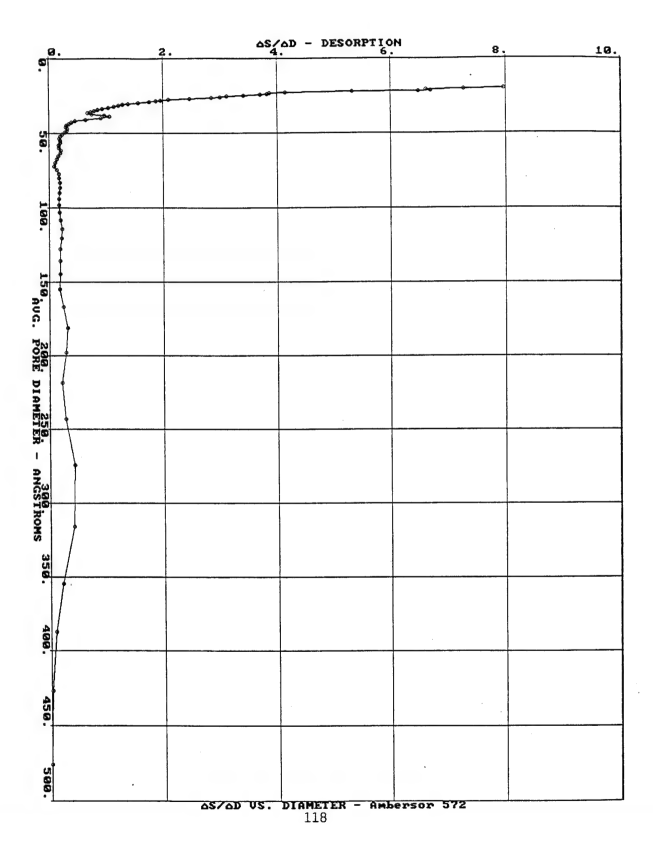
Standard Deviation = 74.3549 Angstroms

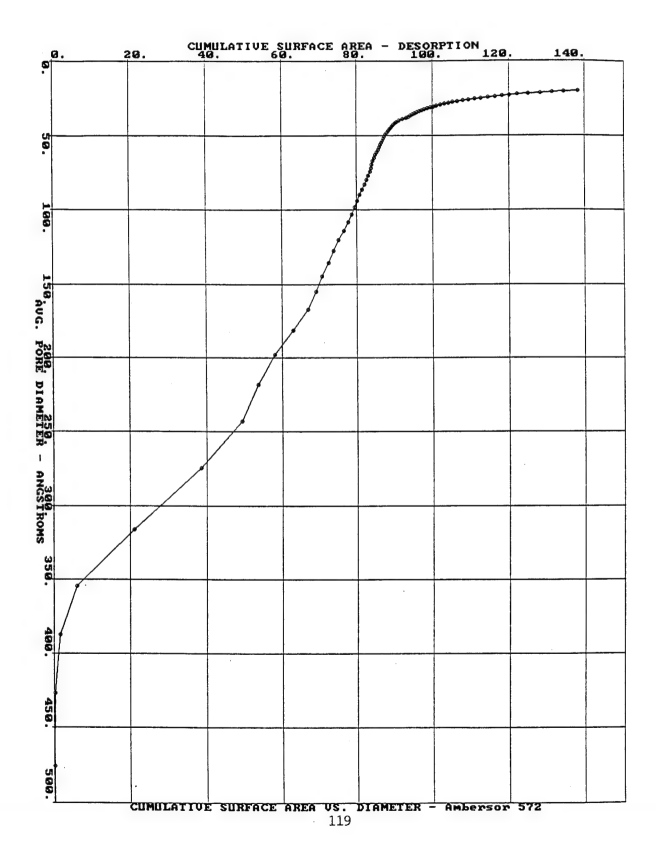
Median Pore Diameter (based on surface area) = 154.502 Angstroms

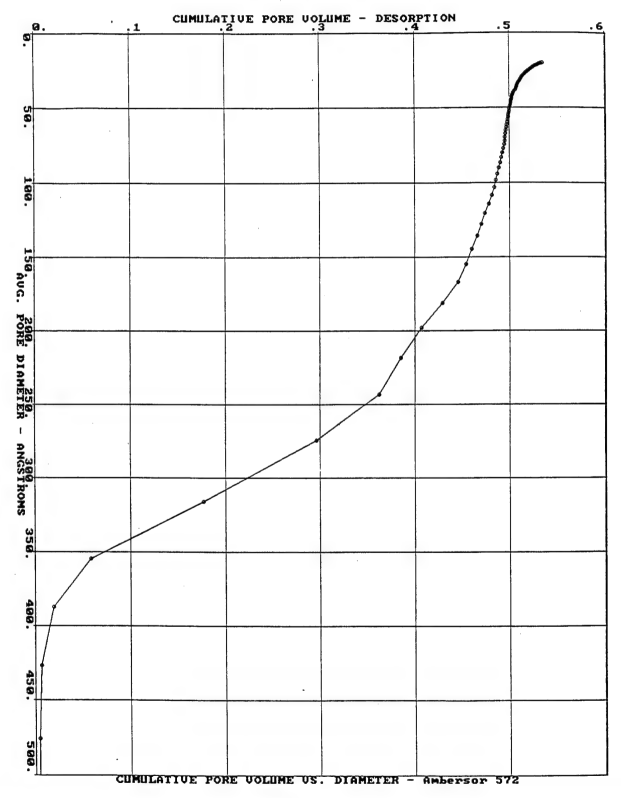
Standard Deviation = 28.0087 Angstroms











#### BET SURFACE AREA ANALYSIS

### 12-16-1994

POROUS MATERIALS, INC. ANALYTICAL SERVICES DIVISION CORNELL INDUSTRY RESEARCH PARK, BLDG. 4 ITHACA, NY 14850 USA PHONE (607) 257-4267 OR 257-5544

NOTE: RESULTS CALCULATED USING WEIGHT AFTER OUTGASSING. TO GET RESULTS USING ORIGINAL WEIGHT, MULTIPLY THESE VALUES BY WT AFTER OUTGAS/WT BEFORE OUTGAS

MTU

Ed. Marchand

SAMPLE ID: Amsorcat (MTU-CAT)

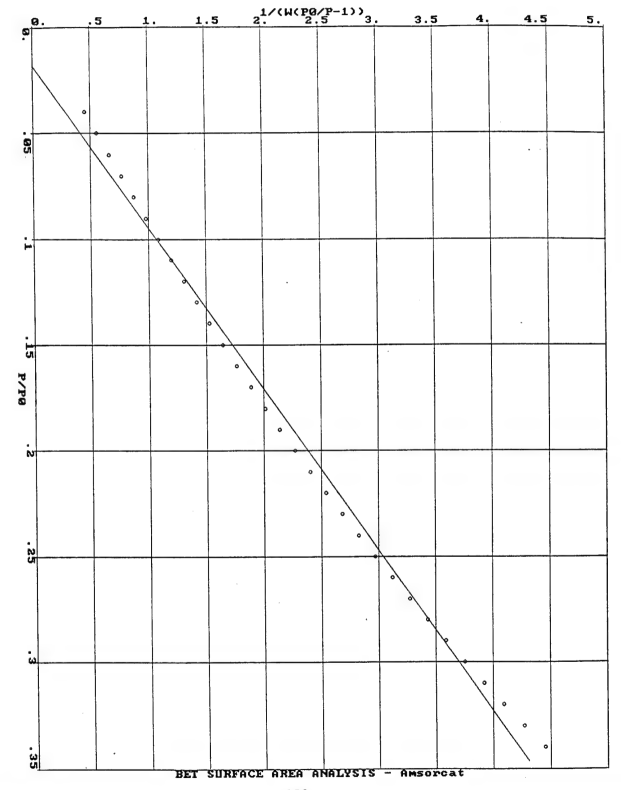
SPECIFIC SURFACE AREA = 324.7001 M2/GM

.8409 SAMPLE WEIGHT= GM BEFORE OUTGASSING .831 GM AFTER OUTGASSING Sample Density= 1.5 BET C VALUE= -52.112 SLOPE= 13.161 GM-1 Y INTERCEPT= -.248 GM-1 CORRELATION COEFFICIENT= .996 ADSORBATE= NITROGEN VA VOLUME= 44.9502 CC Vit VOLUME= 11.14068 CC Vln VOLUME= 14.155 CC INSTRUMENT TEMPERATURE= 305.36 ROOM TEMPERATURE= 296.7 K W = Wt. of adsorbed gas per gram of sample

SAMPLE OUTGASSED AT 20 DEGREES CELSIUS

	P	P0 -1
DATA	-	(W(1))
POINT	P0	P
9	0.04000000	0.44900000
10	0.05000000	0.55200000
11	0.0600000	0.65900000
12	0.0700000	0.76600000
13	0.0800000	0.87300000
14	0.09000001	0.97900000
15	0.1000000	1.085
16	0.11000000	1.194
17	0.12000000	1.304
18	0.13000000	1.415
19	0.14000000	1.526
20	0.15000000	1.641
21	0.16000000	1.762
22	0.17000000	1.885
23	,0.18000000	2.011
24	0.19000000	2.138

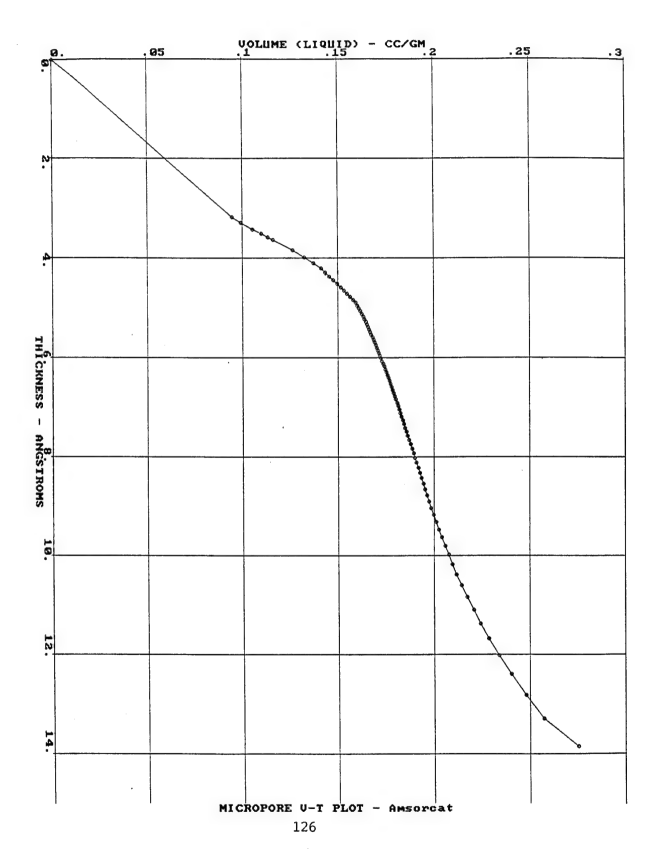
25	0.2000000	2.268
26	0.21000000	2.402
27	0.22000000	2.54
28	0.2300000	2.68
29	0.2400000	2.824
30	0.25000000	2.971
31	0.2600000	3.12
32	0.2700000	3.273
33	0.2800000	3.429
34	0.2900000	3.588
35	0.3000000	3.752
36	0.31000000	3.92
37	0.3200000	4.093
38	0.33000000	4.27
39	0.3400000	4.453



## MICRO PORE CALCULATIONS

Thickness Angstr		Volume (Liquid) cc/g	Hydraulic Radius Angstroms
Angstrands	3.425313.3.6384.19989564.1330.3.3.3.3.3.3.3.3.3.3.3.3.3.3.3.3.3.	0.095 0.099 0.105	
6.586 - 6.648 - 6.711 -	6.648 6.711 6.775	0.178 0.179 0.179	6.617 6.679 6.743

6.840	-	6.906	0.180	6.873
6.906	_	6.974	0.181	6.940
6.974	-	7.043	0.181	7.009
7.043	-	7.114	0.182	7.078
7.114	-	7.186	0.183	7.150
7.186	-	7.259	0.183	7.222
7.259	-	7.335	0.184	7.297
7.335	-	7.412	0.184	7.373
7.412	-	7.491	0.185	7.452
7.491	-	7.572	0.186	7.532
7.572	-	7.656	0.186	7.614
7.656	-	7.742	0.187	7.699
7.742	_	7.830	0.188	7.786
7.830	_	7.921	0.189	7.876
7.921	_	8.015	0.189	7.968
8.015	_	8.112	0.190	8.064
8.112	_	8.212	0.191	8.162
8.212	_	8.316	0.192	8.264
8.316	_	8.424	0.193	8.370
8.424	_	8.536	0.193	8.480
8.536	_	8.652	0.194	8.594
8.652	_	8.773	0.195	8.712
8.773	_	8.899	0.196	8.836
8.899	_	9.031	0.197	8.965
9.031	_	9.170	0.199	9.101
9.170	_	9.315	0.200	9.242
9.315	_	9.468	0.201	9.391
9.468	-	9.629	0.202	9.548
9.629	_	9.799	0.204	9.714
9.799	_	9.980	0.206	9.890
9.980	-	10.172	0.208	10.076
10.172		10.378	0.210	10.275
10.378	_	10.598	0.212	10.488
10.598	_	10.836	0.214	10.717
10.836	-	11.092	0.217	10.964
11.092	_	11.372	0.221	11.232
11.372	-	11.678	0.224	11.525
11.678	-	12.017	0.228	11.847
12.017	-	12.393	0.234	12.205
12.393		12.816	0.240	12.605
12.816	-	13.299	0.248	13.058
13.299	-	13.856	0.257	13.577



### T-PLOT ANALYSIS:

T-PLOT INTERCEPT:

0.10706

T-PLOT SLOPE:

0.010190

MICROPORE VOLUME:

0.10706 CC/GM

FILM THICKNESS USED BETWEEN 8.000 and 12.000 ANGSTROMS

SURFACE AREA = 101.8956 SQ. M/GM (MESO AND MACROPORES ONLY)

MICROPORE SURFACE AREA = 222.8045 SQ. M/GM

## GAS ADSORPTION/DESORPTION RESULTS

ADSORBATE: NITROGEN

## ISOTHERM AND PORE VOLUME DISTRIBUTION

## ADSORPTION

								•	
	PORE	VOLUME	LIQ. VOL	AVE.	CUMUL.	Δv	∆s	SURFACE	CUMUL.
P/P0	DIAMETER	ADSORBED	ADSORBED	POR.DIAM	POR. VOL			AREA S	. AREA
	Å	CC/GM STP	CC/GM	Å	CC/GM	ΔD	ΔD	M2/GM	M2/GM
1.000	141650	340.1398	0.52382	141655	-0-	8.00E-08	4.37E-08	-0-	-0-
0.997	5711.8	333.0673	0.51292	73683.3	0.01094	3.99E-06	3.32E-05	0.00594	0.00594
0.995	3883.5	328.4128	0.50576	4797.63	0.01833	6.95E-06	8.40E-05	0.06167	0.06761
0.990	1957.7	316.3942	0.48725	2920.59	0.03762	1.92E-05	4.45E-04	0.26416	0.33177
0.985	1313.5	307.5428	0.47362	1635.60	0.05205	4.44E-05	0.00146	0.35277	0.68453
0.980	990.44	296.7394	0.45698	1151.97	0.06995	7.85E-05	0.00331	0.62173	1.30626
0.975	796.1	286.0157	0.44046	893.270	0.08794	1.22E-04	0.00630	0.80550	2.11176
0.970	666.21	275.1239	0.42369	731.155	0.10642	1.77E-04	0.01080	1.01099	3.12275
0.965	573.22	264.2112	0.40689	619.717	0.12513	2.40E-04	0.01695	1.20758	4.33033
0.960	503.33	253.3761	0.39020	538.275	0.14389	3.12E-04	0.02505	1.39380	5.72413
0.955	448.85	242.5921	0.37359	476.088	0.16273	3.99E-04	0.03585	1.58299	7.30712
0.950	405.18	231.6586	0.35675	427.013	0.18201	4.95E-04	0.04920	1.80617	9.1133
0.945	369.37	220.7989	0.34003	387.274	0.20132	5.85E-04	0.06400	1.99469	11.1080
0.940	339.48	210.2879	0.32384	354.426	0.22015	6.40E-04	0.07550	2.12490	13.2329
0.930	292.37	193.0715	0.29733	315.927	0.25120	6.95E-04	0.09600	3.93120	17.1641
0.920	256.92	178.8804	0.27548	274.646	0.27698	7.55E-04	0.11850	3.75433	20.9184
0.910	229.24	167.036	0.25724	243.079	0.29861	6.05E-04	0.10450	3.56084	24.4792
0.900	207.02	161.056	0.24803	218.133	0.30898	4.69E-04	0.09050	1.90000	26.3792
0.890	188.79	156.0846	0.24037	197.905	0.31756	4.78E-04	0.10150	1.73565	28.1149
0.880	173.54	151.8067	0.23378	181.161	0.32495	4.67E-04	0.10750	1.63075	29.7456
0.870	160.59	148.3571	0.22847	167.063	0.33080	4.18E-04	0.10350	1.40012	31.1458
0.860	149.46	145.7035	0.22438	155.025	0.33512	3.88E-04	0.10350	1.11521	32.2610
0.850	139.78	143.3818	0.22081	144.619	0.33887	4.18E-04	0.11950	1.03637	33.2973
0.840	131.28	141.1017	0.21730	135.532	0.34265	4.29E-04	0.13050	1.11570	34.4130
0.830	123.76	139.1696	0.21432	127.524	0.34577	3.92E-04	0.12650	0.97837	35.3914
0.820	117.06	137.5602	0.21184	120.412	. 0.34826	3.50E-04	0.11950	0.82750	36.2189
0.810	111.04	136.2071	0.20976	114.050	0.35025	3.50E-04	0.12600	0.69903	36.9179
0.800	105.61	134.8885	0.20773	108.323	0.35224	3.76E-04	0.14250	0.73594	37.6539
0.790	100.67	133.6413	0.20581	103.141	0.35414	3.78E-04	0.15000	0.73619	38.3901
0.780	96.177	132.5179	0.20408	98.4258	0.35581	3.68E-04	0.15300	0.67868	39.0687
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0.00200		64.38936							
0.00101		61.47793							

### SUMMARY SHEET

MTU

Ed. Marchand

SAMPLE ID: Amsorcat

Total Surface Area = 324.7001 sq. m/gm

Average Pore Diameter (4V/S) = 64.8157 Angstroms

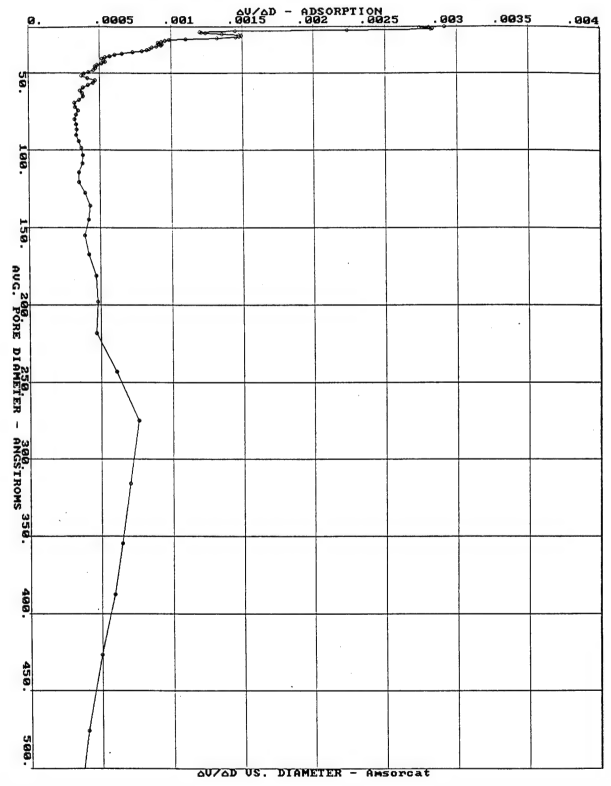
Total Pore Volume = 0.5261 cc/g

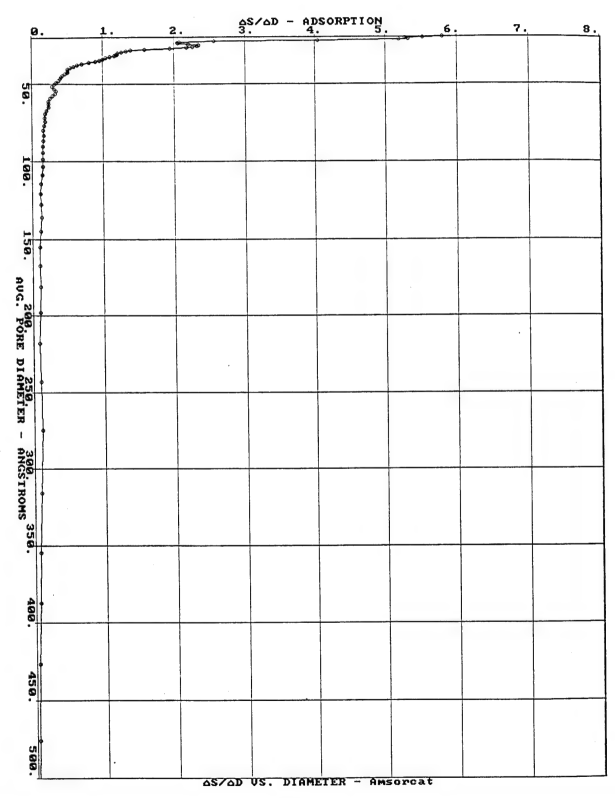
Median Pore Diameter (based on pore volume) = 296.913 Angstroms

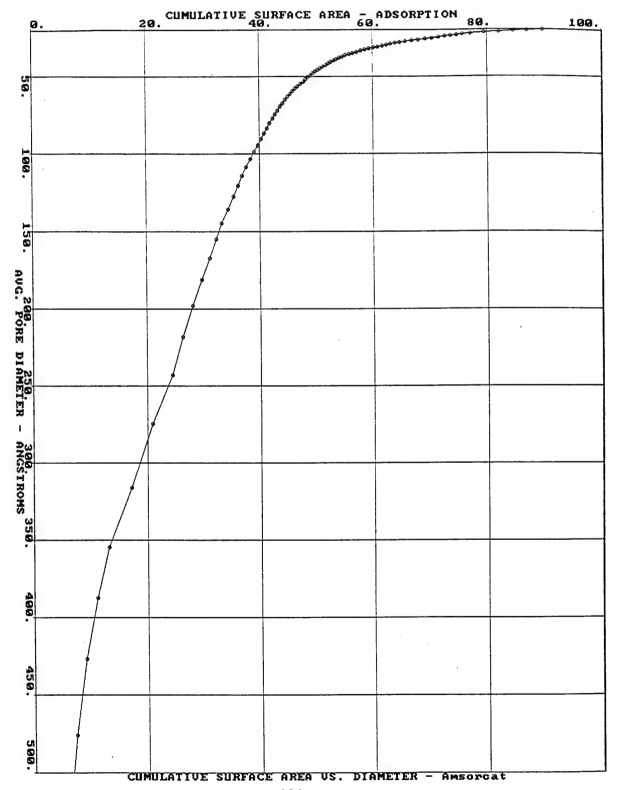
Standard Deviation = 79.3122 Angstroms

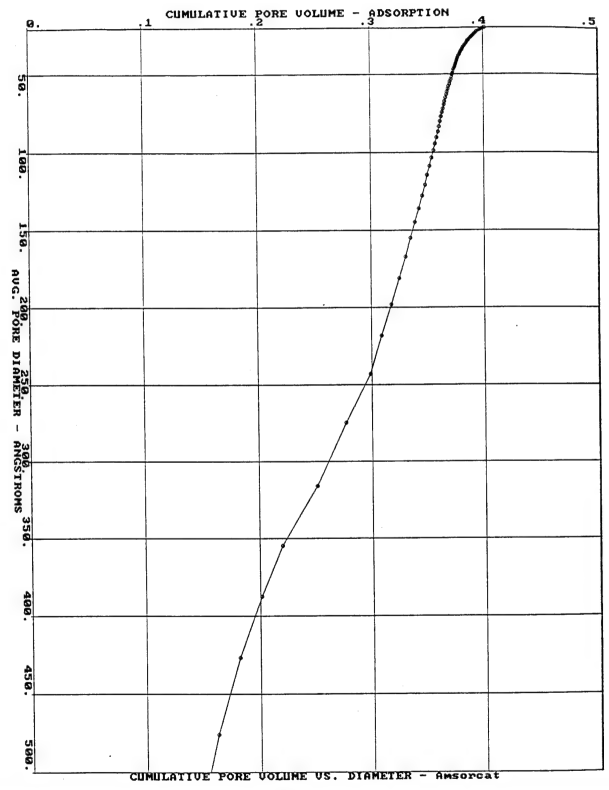
Median Pore Diameter (based on surface area) = 63.974 Angstroms

Standard Deviation = 23.5888 Angstroms









# GAS ADSORPTION/DESORPTION RESULTS

## ADSORBATE: NITROGEN

# ISOTHERM AND PORE VOLUME DISTRIBUTION

## DESORPTION

	PORE	VOLUME	LIQ. VOL	AVE.	CUMUL.	Λv	∆s	SURFACE	CUMUL.
P/P0	DIAMETER	ADSORBED	ADSORBED	POR.DIAM	POR. VOL				. AREA
	Å	CC/GM STP	CC/GM	Å	CC/GM	ΔD	Λp	M2/GM	M2/GM
0.995	3883.5	339.3051	0.52253	3883.49	-0-		9.65E-06	-0-	-0-
0.990	1957.7	338.4578	0.52122	2920.59	0.00136	1.35E-06	3.11E-05	0.01867	0.01867
0.985	1313.5	337.841	0.52028	1635.60	0.00237	2.41E-06	7.80E-05	0.02462	0.04330
0.980	990.44	337.2859	0.51942	1151.97		2.61E-06		0.03188	0.07517
0.975	796.1	336.9962	0.51897	893.270	0.00377	2.31E-06	1.17E-04	0.02144	0.09662
0.970	666.21	336.8216	0.51871	731.155	0.00405	2.61E-06	1.59E-04	0.01573	0.11235
0.965	573.22	336.6602	0.51846	619.717		4.65E-06		0.01738	0.12973
0.960	503.33	336.4172	0.51808	538.275	0.00474	1.54E-05	0.00127	0.03103	0.16076
0.955	448.85	335.7192	0.51701	476.088	0.00598	5.55E-05	0.00510	0.10388	0.26464
0.950	405.18	333.7324	0.51395	427.013	0.00956	2.11E-04	0.02140	0.33520	0.59984
0.945	369.37	327.5268	0.50439	387.274	0.02088	6.55E-04	0.07250	1.16943	1.76927
0.940	339.48	312.2777	0.48091	354.426	0.04899	0.00125	0.15050	3.17304	4.94231
0.930	292.37	268.3468	0.41325		0.13101	0.00160	0.21800	10.3837	15.3260
0.920	256.92	240.1876	0.36989	274.646	0.18369	0.00127	0.19700	7.67239	22.9984
0.910	229.24	223.6478	0.34442	243.079	0.21434	0.00125	0.22050	5.04465	28.0430
0.900	207.02	207.4409	0.31946	218.133	0.24481	0.00165	0.32050	5.58637	33.6294
0.890	188.79	189.6526	0.29207	197.905	0.27891	0.00229	0.48900	6.89328	40.5227
0.880	173.54	169.1	0.26041	181.161	0.31915	0.00221	0.50500	8.88455	49.4072
0.870	160.59	156.6159	0.24119	167.063	0.34299	0.00118	0.28800	5.70731	55.1145
0.860	149.46	152.3343	0.23459	155.025	0.34976	6.10E-04	0.16450	1.74638	56.8609
0.850	139.78	148.5173	0.22872	144.619	0.35577	6.05E-04	0.17450	1.66241	58.5233
0.840	131.28	145.2164	0.22363	135.532	0.36086	4.98E-04	0.15050	1.50386	60.0272
0.830	123.76	142.9145	0.22009	127.524	0.36392	4.18E-04	0.13500	0.96099	60.9882
0.820	117.06	140.7715	0.21679	120.412	0.36679	3.93E-04	0.13350	0.95306	61.9413
0.810	111.04	138.9967	0.21405	114.050	0.36897	3.36E-04	0.12100	0.76212	62.7034
0.800	105.61	137.4786	0.21172	108.323	0.37067	3.39E-04	0.12850	0.63068	63.3341
0.790	100.67	135.9785	0.20941	103.141	0.37246	3.36E-04	0.13350	0.69180	64.0259
0.780	96.177	134.6802	0.20741	98.4258	0.37387	2.72E-04	0.11250	0.57298	64.5988
0.770	92.057	133.6074	0.20576	94.1170	0.37483	2.52E-04	0.10950	0.40883	65.0077
0.760	88.269	132.5473	0.20412	90.1629	0.37585	2.83E-04	0.12800	0.45259	65.4603
0.750	84.77201	131.516	0.20253	86.5203	0.37688	2.45E-04	0.11500	0.47675	65.9370

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                    120.4081
0.520
                              0.18491 42.6846 0.38095 1.49E-04 0.14150 0.12476 68.2430
0.510
          42.196 . 120.0693
                              0.18438 41.7237 0.38111 1.87E-04 0.18200 0.14655 68.3896
0.500
          41.252
                    119.7304
                              0.18385 40.7962 0.38130 2.40E-04 0.23800 0.18945 68.5790 0.18331 39.9003 0.38153 2.82E-04 0.28600 0.23567 68.8147
                    119.3852
0.490
          40.341
0.480
          39.46
                    119.0331
                               0.18277 39.0339 0.38179 3.12E-04 0.32300 0.25912 69.0738
0.470
          38.608
                    118.6807
                              0.18223 38.1954 0.38206 3.37E-04 0.35650 0.28153 69.3554
          37.783
                    118.3283
0.460
                               0.18169 37.3833 0.38233 3.15E-04 0.34050 0.29720 69.6526
                    117.9776
 0.450
          36.984
                               0.18118 36.5959 0.38255 2.67E-04 0.29400 0.24083 69.8934
 0.440
          36.208
                    117.6463
                                                0.38274 2.61E-04 0.29400 0.20948 70.1029
                               0.18068 35.8319
          35.456
                    117.3274
 0.430
                               0.18019 35.0901 0.38294 2.84E-04 0.32750 0.22675 70.3296
 0.420
          34.725
                    117.009
                               0.17970 34.3691 0.38315 2.98E-04 0.35000 0.24506 70.5747
                    116.6908
 0.410
          34.014
                               0.17922 33.6679 0.38336 2.88E-04 0.34600 0.24597
                                                                                    70.8206
 0.400
          33.322
                    116.3765
                               0.17875 32.9853 0.38355 2.86E-04 0.35000 0.22693
                                                                                    71.0476
 0.390
          32.648
                    116.07
                               0.17828 32.3203 0.38374 3.05E-04 0.38200 0.23841 71.2860
          31.992
                    115.7647
 0.380
                               0.17781 31.6720 0.38394 3.28E-04 0.41800 0.25642 71.5424
                    115.4592
 0.370
          31.352
                               0.17734 31.0394 0.38415 3.54E-04 0.46050 0.27243 71.8148
 0.360
          30.727
                    115.1538
                               0.17686 30.4217 0.38438 4.07E-04 0.54000 0.29613 72.1110
                    114.847
 0.350
          30.116
                               0.17638 29.8181 0.38464 4.64E-04 0.62500
                                                                           0.35552
                                                                                    72.4665
                    114.5325
 0.340
          29.52
                                                                           0.38694 72.8534
                                                0.38493 4.95E-04 0.68000
 0.330
          28.936
                    114.2153
                               0.17589 29.2277
                                                0.38521 5.10E-04 0.72500 0.40306 73.2565
          28.364
                    113.8979
                               0.17540 28.6498
 0.320
                               0.17491 28.0837 0.38551 5.55E-04 0.80500 0.41773 73.6742
                    113.5804
 0.310
           27.804
                               0.17441 27.5287 0.38583 6.80E-04 1.00500 0.47511 74.1493
                    113.2561
 0.300
           27.254
                               0.17389 26.9841 0.38625 7.90E-04 1.18500 0.61696 74.7663
           26.714
                    112.9127
 0.290
                               0.17335 26.4492 0.38668 8.15E-04 1.25000 0.65178
                                                                                    75.4181
 0.280
           26.184
                    112.5661
                               0.17282 25.9234 0.38711 8.25E-04 1.29000 0.66278 76.0809
           25.663
                    112.2195
 0.270
                                                0.38754 8.65E-04 1.37500 0.67311 76.7540
                               0.17228 25.4061
 0.260
           25.149
                     111.8727
                               0.17174 24.8966 0.38799 0.00096 1.55000 0.72919 77.4832
 0.250
           24.644
                     111.5196
                               0.17118 24.3943 0.38850 0.00102 1.68500 0.82953 78.3127
                    111.1547
           24.145
 0.240
                               0.17061 23.8985 0.38900 0.00102 1.72000 0.84512
                                                                                    79.1578
 0.230
           23.652
                    110.7882
                               0.17005 23.4087 0.38950 0.00100 1.73500 0.84425 80.0021 0.16948 22.9242 0.38998 0.00105 1.84000 0.83712 80.8392
 0.220
           23.165
                     110.4216
                     110.0551
 0.210
           22.683
                                        22.4443 0.39050 0.00134 2.41000 0.93188 81.7711
                                0.16890
 0.200
           22.205
                     109.6774
                                0.16825 21.9683 0.39125 0.00163 3.00500 1.36002 83.1311
 0.190
           21.731
                     109.2547
                                0.16758 21.4954 0.39204 0.00166 3.11500 1.48210 84.6132
           21.26
                     108.818
 0.180
                                0.16691 21.0250 0.39281 0.00159 3.06000 1.45147
                                                                                    86.0647
 0.170
           20.79
                     108.3812
                                0.16623 20.5561 0.39354 0.00188 3.70500 1.41988
                                                                                    87.4845
 0.160
           20.322
                     107.9433
                               0.16547 20.0878 0.39457 0.00220 4.38000 2.05188 89.5364
           19.854
                     107.4467
 0.150
```

0.140	106.8453
0.130	106.2304
0.120	105.6154
0.110	104.9997
0.100	104.2879
0.0900	103.2498
0.0800	102.1306
0.0700	101.0109
0.0600	99.88666
0.0500	98.40459
0.0400	95.953
0.0300	93.34013
0.0200	90.05223
0.0100	83.99156
0.00800	82.30693
0.00600	80.3519
0.00400	77.44178
0.00200	73.68878
0.00100	70 93205

### SUMMARY SHEET

MTU

Ed. Marchand

SAMPLE ID: Amsorcat

Total Surface Area = 324.7001 sq. m/gm

Average Pore Diameter (4V/S) = 64.6566 Angstroms

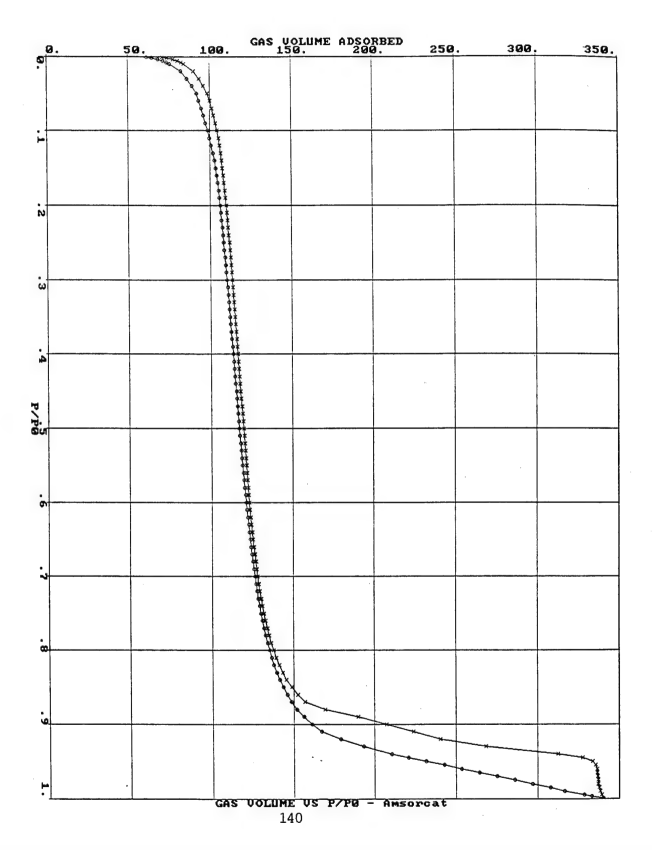
Total Pore Volume = 0.5249 cc/g

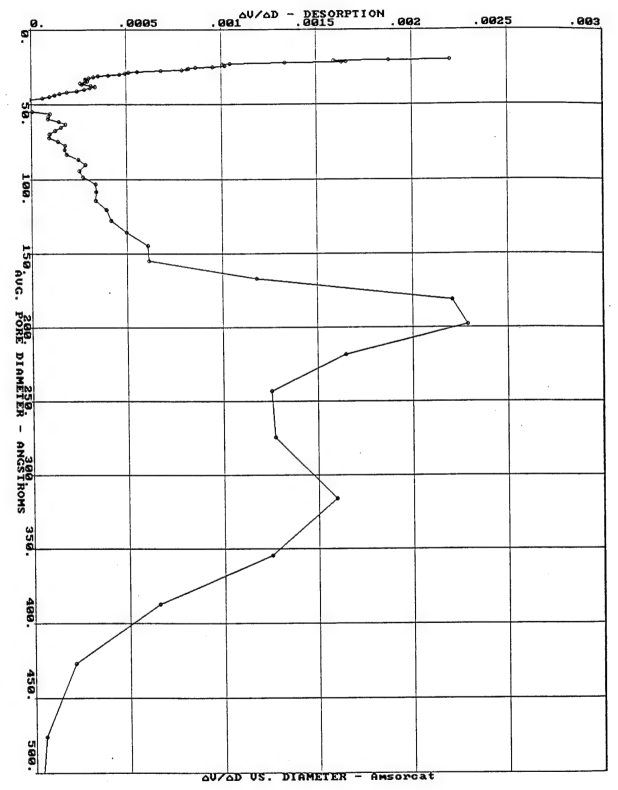
Median Pore Diameter (based on pore volume) = 207.682 Angstroms

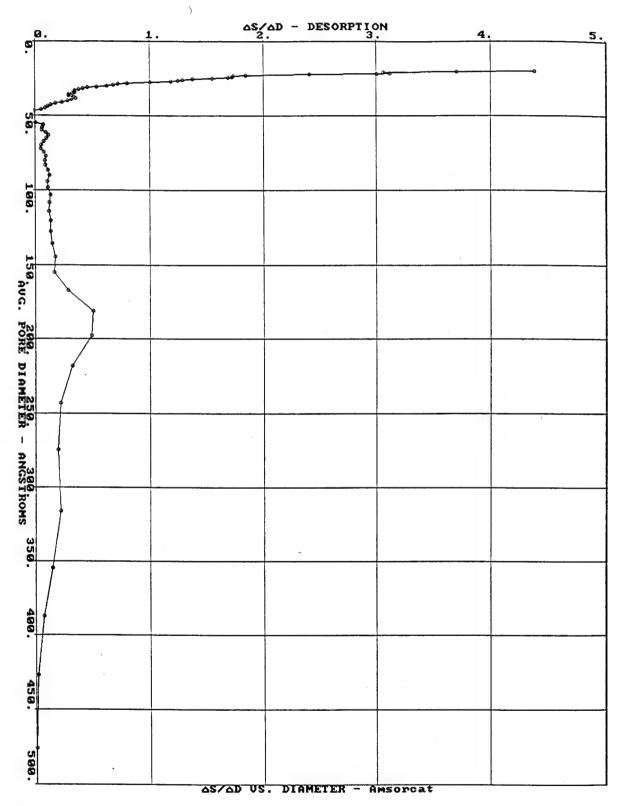
Standard Deviation = 86.4755 Angstroms

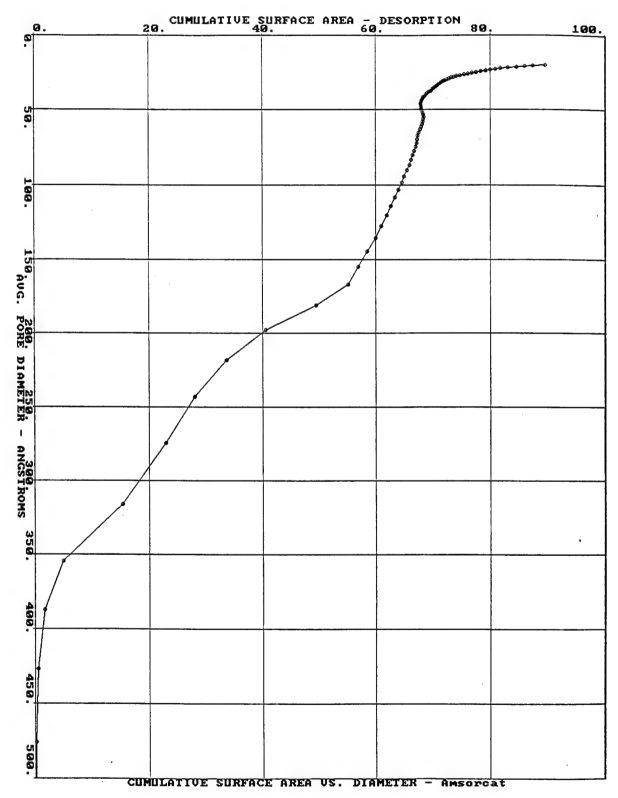
Median Pore Diameter (based on surface area) = 189.903 Angstroms

Standard Deviation = 36.1261 Angstroms









# APPENDIX B

Quattro Pro Spreadsheets with Data for the Figures Used in this Dissertation

Gas adsorption/desorption results from PMI, Inc. (Figure 2-3, PhD Dissertation, 2/96)
Analysis done 16 Dec 1994

P/Po	Po ADSORPTION		DESORPTION	
	MTU-CAT AM 572		MTU-CAT AM 572	
1	340			
0.997	333			
0.995	328	586	339	
0.99	316	585	338.4	585.5
0.985	308	583	337.8	584.7
0.98	297	579	337.3	584.1
0.975	286	567	337	583.9
0.97	275	546	336.8	583.6
0.965	264	521	336.7	583.4
0.96	253	497	336.4	583.1
0.955	242	474	335.7	582.7
0.95	232	454	333.7	581.7
0.945	221	438	327.5	574.7
0.94	210	424	312.3	553.6
0.93	193	398	268.3	490.3
0.92	179	378	240.2	426.9
0.91	167	366	223.6	391.9
0.9	161	355	207.4	378
0.89	156	348	189.6	366
0.88	151.8	342	169.1	354
0.87	148.4	338	156	344.8 339.5
0.86	145.7	333.8	152.3	
0.85	143.4	330.2	148.5	335.4
0.84	141.1	327.2	145.2	331.6
0.83	139.2	324.8	142.9	328.6 325.8
0.82	137.6	322.6	140.8 139	323.6
0.81 0.8	136.2 134.9	320.6 318.8	137.5	320.6
0.8	134.9	317.2	137.5	318.7
0.78	132.5	317.2	134.7	317
0.75	131.5	314.2	133.6	315.5
0.76	130.6	312.8		314.1
0.75	129.7	311.6	131.5	312.8
0.74	128.9	310.5	130.7	311.5
0.73	128.2	309.5	129.9	310.4
0.72	127.5	308.5	129.2	309.4
0.71	126.8	307.7	128.4	308.4
0.7	126.2	307	127.8	307.6
0.69	125.6	306.2	127.3	306.8
0.68	125	305.5	126.7	306.1
0.67	124.5	304.8	126.2	305.4
0.66	123.9	304.1	125.6	304.6
0.65	123.4	303.4	125.1	303.9
0.64	122.9	302.8	124.6	303.2
0.63	122.4	302.2	124.2	302.5
0.62	121.9	301.6	123.7	301.9

0.61	121.4	300.9	123.3	301.3
0.6	120.9	300.3	122.9	300.7
0.59	120.5	299.7	122.6	300.1
0.58	120.1	299.2	122.3	299.6
0.57	119.7	298.7	122	299
0.56	119.3	298.1	121.7	298.4
0.55	118.9	297.6	121.4	297.9
0.54	118.5	297.1	121.1	297.4
0.53	118.1	296.5	120.7	296.8
0.52	117.8	296	120.4	296.3
0.51	117.4	295.5	120.1	295.8
0.5	117	295	119.7	295.3
0.49	116.6	294.5	119.4	294.8
0.48	116.3	293.9	119	294.2
0.47	116	293.4	118.7	293.6
0.46	115.6	292.9	118.3	293
0.45	115.2	292.4	118	292.5
	114.9	291.9	117.6	292
0.44				291.5
0.43	114.5	291.4	117.3	
0.42	114.2	290.9	117	291 290.5
0.41	113.8	290.4	116.7	
0.4	113.4	289.9	116.4	290
0.39	113.1	289.4	116.1	289.5
0.38	112.7	288.9	115.8	289
0.37	112.3	288.4	115.4	288.5
0.36	112	287.9	115.2	288
0.35	111.6	287.4	114.8	287.5
0.34	111.3	286.8	114.5	287
0.33	111	286.3	114.2	286.5
0.32	110.6	285.7	113.9	286
0.31	110.3	285.2	113.6	285.5
0.3	109.9	284.6	113.2	285
0.29	109.5	284.1	112.9	284.4
0.28	109.1	283.6	112.6	283.8
0.27	108.7	283	112.2	283.3
0.26	108.3	282.5	111.9	282.7
0.25	107.9	281.9	111.5	282.1
0.24	107.6	281.3	111.2	281.5
0.23	107.2	280.8	110.8	280.9
0.22	106.8	280.2	110.4	280.3
0.21	106.5	279.5	110	279.7
0.2	106	278.8	109.7	279
0.19	105.5	278	109.2	278.3
0.18	105	277.3	108.8	277.6
0.17	104.5	276.6	108.4	276.8
0.16	104	275.8	107.9	276.1
0.15	103.4	274.9	107.4	275.2
0.14	102.6	273.9	106.8	274.2
0.13	101.6	272.9	106.2	273.2
0.12	100.6	271.9	105.6	272.2
0.11	99.5	270.6	105	271.1
0.1	98.5	268.8	104.3	269.9
V. I	30.0	200.0	10-710	

0.09	97.2	266.8	103.2	268
0.08	95.8	264.9	102.1	265.9
0.07	94.5	263	101	263.8
0.06	93.1	261	99.9	261.8
0.05	91.6	258.3	98.4	259
0.04	89.1	253.5	96	254
0.03	85.9	248	93.3	248.5
0.02	82	241	90	241.6
0.01	75.3	227	84	228
0.008	73.5	223	82.3	224.6
0.006	71.4	218	80.4	220
0.004	68.3	211	77.4	214
0.002	64.4	202	73.7	206
0.00101	61.5	195	70.8	199.9

Figure 2-4, PhD Dissertation, 2/96 BET Adsorption Isotherm and Pore Volume Distribution data from PMI, Inc.

Plain Amb	ersorb 57	2	SuperCat
Pore	^ V/ ^ D	^S/^D	^V/^D ^S/^D
Dia (A)			
1958	2E-06	5 <b>E-0</b> 5	1.4E-06 3.1E-05
1313	2E-06	8 <b>E-0</b> 5	2.4E-06 7.8E-05
990	2E-06	0.0001	2.6E-06 0.00011
796	3E-06	0.00016	2.3E-06 0.00012
666	4E-06	0.00025	2.6E-06 0.00016
573	6E-06	0.0004	4.7E-06 0.00033
503	9E-06	0.0007	1.5E-05 0.00127
448	3E-05	0.0026	5.6E-05 0.0051
405	0.0002	0.0221	0.00021 0.0214
369	0.00087	0.097	0.00066 0.0725
339	0.00177	0.213	0.00125 0.1505
292	0.003	0.416	0.0016 0.218
256	0.0028	0.434	0.00127 0.197
229	0.0016	0.28	0.00125 0.22
207	0.0011	0.22	0.00165 0.32
188	0.0013	0.285	0.00229 0.489
173	0.0014	0.312	0.00221 0.505
160	0.001	0.247	0.00118 0.288
149	0.0007	0.183	0.00061 0.1645
139	0.00065	0.186	0.0006 0.1745
131	0.0006	0.187	0.0005 0.1505
123	0.0006	0.188	0.00042 0.135
117	0.0006	0.22	0.00039 0.1335
111	0.0006	0.23	0.00034 0.121
105	0.0005	0.203	0.00034 0.128
100	0.0005	0.185	0.00034 0.133
96	0.0004	0.17	0.00027 0.112
92	0.0004	0.1705	0.00025 0.109
88	0.0004	0.18	0.00028 0.128
85	0.0004	0.189	0.00025 0.115
81	0.0004	0.1875	0.00019 0.09
78	0.00034	0.174	0.00017 0.088

76	0.00033	0.176	0.00018	0.094
73	0.00026	0.139	0.00014	0.076
71	0.00017	0.097	9E-05	0.052
68	0.00018	0.106	9.8 <b>E-</b> 05	0.058
66	0.00022	0.134	0.00013	0.076
64	0.00026	0.164	0.00016	0.098
62	0.00031	0.197	0.00018	0.116
60	0.00031	0.206	0.00014	0.095
59	0.00026	0.176	9E-05	0.061
57	0.00024	0.169	9E-05	0.064
55	0.00027	0.197	0.0001	0.072
54	0.00027	0.198	7.7E-06	0.0049
52.6	0.00024	0.185	-0.0001	-0.078
51	0.00026	0.201	-0.0001	-0.093
50	0.0003	0.237	-0.0001	-0.0825
49	0.00034	0.277	-8.6E-05	-0.0705
47	0.00038	0.317	-6.3E-05	-0.053
46	0.00037	0.322	-6.3E-06	-0.0049
45	0.00034	0.296	6 <b>E</b> -05	0.054
44	0.00034	0.304	9.8E-05	0.089
43	0.00038	0.347	0.00012	0.114
42	0.00041	0.393	0.00015	0.141
41	0.00047	0.455	0.00019	0.182
40	0.00064	0.64	0.00024	0.238
39.5	0.00089	0.905	0.00028	0.286
38.6	0.001	1.06	0.00031	0.323
37.8	0.00092	0.97	0.00034	0.356
37	0.0007	0.75	0.00032	0.34
36	0.00061	0.675	0.00027	0.294
35.4	0.00064	0.73	0.00026	0.294
34.7	0.00069	0.795	0.00028	0.327
34	0.00073	0.86	0.0003	0.35
33.3	0.00078	0.935	0.00029	0.346
32.6	0.00085	1.04	0.00029	0.35
32	0.00091	1.135	0.00031	0.382
31.4	0.00095	1.215	0.00033	0.418
30.7	0.00099	1.295	0.00035	0.46
30.1	0.001	1.385	0.00041	0.54
29.5	0.00115	1.56	0.00046	0.625
29	0.00127	1.76	0.0005	0.68

28.4	0.00133	1.875	0.00051	0.725
27.8	0.00136	1.96	0.00056	0.805
27.2	0.00143	2.1	0.00068	1.005
26.7	0.00165	2.475	0.00079	1.185
26.2	0.00187	2.85	0.00082	1.25
25.7	0.00193	3	0.00083	1.29
25.1	0.00196	3.12	0.00087	1.38
24.6	0.0021	3.41	0.00096	1.55
24.1	0.00224	3.705	0.00102	1.68
23.6	0.00226	3.82	0.00102	1.72
23.1	0.00224	3.86	0.001	1.74
22.7	0.00235	4.14	0.00105	1.84
22.2	0.00295	5.3	0.00134	2.41
21.7	0.00352	6.45	0.00163	3
21.3	0.00356	6.7	0.00166	3.115
20.8	0.00344	6.6	0.00159	3.06
20.3	0.00369	7.25	0.00188	3.705
20	0.004	7.95	0.0022	4.38

Figure 2-5 in PhD Dissertation 2/96 Huffman Lab results on elemental analysis of catalysts/supports Weight percentages:

•	AM-572-P	MTU-CAT	Methanol rinsed MTU-CA	AT Accuracy (+/- %)
Carbon	95.51	71.11	76.29	0.3
Hydrogen	0.41	0.36	0.92	0.1
Sulfur	1.45	1.31	1.2	0.1
Chlorine	0	21.87	14.29	0.3

Figure 2-6, PhD Dissertation, 2/96

	1320		240379
	TIME(MIN)	GC AREA	PPM EFFLUENT
Jan 2-C	240	1941	5
EBCT=17 s	360	947	. 3
T=190 C	480	645	2
	600	0	0
	720	0	0
	840	0	0
	960	0	0
	1080	0	0
	1200	0	0
	1320	7446	20
	1440	61406	167
	1560	127824	349
	1680	194242	530
	1800	189043	516
	1920	166637	454
	2040	137586	375
	2160	116557	318
	2280	88735	242
•	2400	67220	183
	2520	49764	136
	2640	35321	96
	2760	25618	70
	2880	17368	47
	3000	11559	32
	3120		0
	3240	4935	13
	3360	3593	10
	3480	2572	7
	3600	1950	5
	3720	1471	4
	3840	1141	3
	3960	741	2
	4080	595	2
	4200	381	· 1
	4320	0	0
	4440	0	0

	4560	0		0	
	4680	0		0	
	4800	0		0	
	4920	0		0	
	5040	0		0	
	5160	1073		3	
	5280	7090		19	
	5400	31892		87	
•	5520	72924		199	
	5640	105200		287	
	5760	122000		333	
	5880	125000		341	a.
am572-m	60		1200		3
T=175 C	180		827		2
EBCT=17s	300		680		2
	420		0		0
	540		0		0
	660	•	0		0
	780		0		0
	900		0		0
	1020		0		0
	1140		733		2
	1260		4400		12
	1380		25600		70
	1500		77450		211
	1620		185300		505
	1740		283000		772
	1860		394000		1075
	1980		414000		1129
	2100		401000		1094
	2220		388000		1058
	2340		325000	,	886
	2580		242000		660
•	2820		190000		518
	2940		140000		382
	3060		127000		346
	3660		56000		153
	3780		46000		125
	3900		38600		105

4020	31800	87
4140	27200	74
4260	23500	64
4380	19250	53
4500	15300	42
4620	12100	33
4740	7430	20
4860	5360	15
4980	4124	11
5100	3060	8
5220	2078	6
5340	1684	5
5460	1293	4
5580	1058	3
5700	878	2
5820	690	2
5940	589	2
6060	525	1
6180	0	0
6300	0	0
6420	0	0
6540	0	0
6660	0	0
6780	0	. 0
6900	0	0
7020	0	0
7140	0	0
7260	0	0
7380	0	0
7500	0	0
7620	0	0
7740	552	2
7860	1888	5
7980	7518	21
8100	22269	61
8220	46202	126

Fig 2-8 PhD Dissertation, 2/96 AM 572 plain, 180 C heat tape, then on Joule heating. See page 26/27 of book 6.

				GC area	
Time	Temp	Volts	Ohms	for CCI4	GC Values/10
0	180	0	30	1240	
2.3	180	5	29		
3.3	177	5	26		
4	175	6	24		
5	174	6	28	1541	
6	172	7.5	23		
8	169	9	21		
10	167	9	21		
11	166	12	20		
12	167	12	24	3954	
15.5	166	12	22		
17	165	12.5	21		
20	165	14	19		
21	166	14	22	7094	
29	167	14	22		
31	167	14	. 22		
32	166.5	14	24	6291	
41	165	14.5	21		
46	165	15	22		
59	164	15	27		
61	163	17	26		
63	162	17			
69	160			1977	
70	157	8	11		
71		9	9		
73	158	9	9		
76.2	159			1241	
82.5	160	8	8		
88	165			820	
0					124
5					154
12					395
21					709
32				٨.	629
69					198
76.2					124
88					82

Figure 3-3, PhD Dissertation, 2/96 Temp 38 C unless otherwise noted Cloth Isotherms under various conditions

				Condition
TCE,ppm	15	500	1500	Feed Gas
mg/g = =	18	71	164	Dry
	16	74	131	80% RH, no Joule heating
	15.5	64	129	80% RH, 0.7V (6 mA)
	17			80% RH, 1 V (20 mA)
	1.7			80% RH at 25 C
	7.7			Joule heating capacity @42C
				(still at 80% RH)

Note: RH measured at room temperature (22 C) and not at the temperatures noted above.

Figure 3-4, PhD Dissertation, 2/96 Joulsorption with humidity @ 15 ppmv. Temperature upper, lower, and humidity sens Book 7, pp 93-96, 21 Dec 1995

Adsorbent: K-Cloth, 12 sections in 1/4" glass tube, (0.0167 g) Flow is 48 ml/min at start

Time	T upper	T lower	Humidity @	Temp	Real RH	GC area	Conc (pp
0	00.5	00	F-7	00	50		
5	26.5	28	57 50	26	53		
10	26.5	28	59	27	56		
15	27	28	62	27	58	0	0
20	26.5	28	61	26	55	0	0
30	26.5	27.5	67	26	60	0	
40						44	1610
60						56.6	16.10 20.71
80	26.5	07	00	26	78	50.0	20.71
85 100	26.5	27	82	20	70	69.4	25.39
110	26	27	82	26	78	09.4	25.59
120			84	26	78		
140	26.5	27.5	04	20	70	58.1	21.26
160	25.5	27	83	25	74	40.7	14.89
180	20.0	21	65	20	74	41.7	15.26
200						41.1	15.04
220						39	14.27
240				•		40.7	14.89
260						35.9	13.13
280	24	26	82	23	68	38.2	13.98
284	24.5	26	81	24	73	00.2	. 5.55
288	24.5	26	83	24	74		
292	24.5	26	84	24	75		
296	25	26	86	24	76		
300	25	26	87	25	80	7.85	3.925
304	25		87	24	79		
310	25		88	25	83		
320	25		88	24	80	5.73	2.865
330			85	25	78		
340				25	79	5.11	2.555
350				25	78		
360				25	78	5.85	2.925
370				24	72		
380				25	80	4.7	2.35
390		27	91	25	81		
400						4.1	2.05

420						3.5	1.75
440						3.6	1.8
460						2.72	1.36
480						2.5	1.25
500						2.79	1.395
520						3.03	1.515
540						3.7	1.85
560						5.6	2.8
580						9.7	4.85
600						14	7
620						18.4	9.2
640	,					22.4	11.2
660						24.7	12.35
680						22	11
700						27.8	13.9
720						29.7	14.85
780						31.6	15.8
840						30.8	15.4
900						31	15.5
960						30.8	15.4
1020						29.7	14.85
1080						29.4	14.7
1140						29.1	14.55
1154	24.5	26.5	81	24	71		
1157	24.5	26	80	24	70		
1160	25	26	69	25	66	11.4	5.7
1162	25	26	65	25	60		
1164	24.5	26	64	25	59		
1166	25	26	63	24	55	10.5	5.25
1172	25	26	64	25	59		
1176	25	26	65	25	60		
1180	24.5	26	64	24	57	14.6	7.3
1190	25	26	69	24	61	37.7	18.85
1200	25	26	75	25	71	218.5	109.25
1210	25	26	79	25	77	329.2	164.6
1220	25	26	79	25	77	269.8	134.9
1230	25	26	81	24	73	121.3	60.65
1240	25	26	82	24	73	135.2	67.6
1260	25	26	82	25	78	59.5	29.75
1280	25	26	83	24	74	35.4	17.7
1300						35.1	17.55

Carbon Rod R and R/Ro (McIntos																									0.6316 0.9381	0.6733
R/Ro for packed ACC20	1.00	76.0	0.93	0.90	0.86	0.83	0.79	0.76			•															
3	53	58	27	56	52	24	83	22																		
tion, 2/96 Baudu predictions		0.81	0.796	0.784	0.774	0.766	0.759	0.754	0.751	0.745	0.42	0.45	0.48	0.52	0.57	0.62	0.70	0.78	0.89	0.96	1.00					
Figure 4-3, PhD Dissertation, 2/96 R/Ro for ACC20 and 572	1.00	0.97	0.93	0.90	0.87	0.85	0.81	0.79	0.75	0.72																
Figure 4-3, F R/Ro for AC	91.7	98.6	85.6	82.7	79.9	77.5	74.7	72.1	69.1	66.2	25.6	27.2	29.2	31.6	34.6	37.9	42.5	47.8	54.3	58.4	61					
		9	20	80	06	100	110	120	130	140	150	140	130	120	110	100	83	79	29	61	22	140	123	98	72	20.2

Figure 4-4, PhD Dissertation 2/96 Cloth Adsorbent Isotherms

DRY: (0.0094g cloth)

TIME	GC Area	Conc.pp	Ohm	
0	0		54.2	
8	0		55.13	
15			55.52	
25			55.65	
34			55.76	
41	0			
47			55.76	
61	0			
160			55.79	
181	3.17	2.4		
201	6.65	4.9		
210			56.23	
221	10.35	7.7	55.89	
241	13.84	10.3	56.27	
257			56.2	
271	18.62	13.8	56.76	13 mg TC
281	19.96	14.8	56.67	
361	24.56	18.2	57.07	
421	23.35	17.3	56.88	
541	23.18	17.2	57.05	
661			56.89	
781	20.55	15.3	56.71	
901	20.32	15.1	56.9	
1021	19.61	14.6	56.83	
1141	20.21	15.0	56.93	
1261	18.96	14.1	57.23	
1301	19.57	14.5	56.46	
1321	20.78	15.4	55.78	
1341	22.25	16.5	57.09	
1366	21.24	15.8	55.9	
1381	14.72			
1401	31.29	29.4	55.45	
1421	42.6	40.0	54.82	
1441	94.85	89.1	54.12	
1461	262.72	246.9	54.54	•
1481	388.17	364.8	54.1	

1501	457.82	430.3	54.1	
1521	491.51	461.9	54.18	
1541	458.41	430.8	53.96	
1561	504.05	473.7	54.11	33 mg/g
1581	489.89	460.4	53.82	
1601	484.71	455.6	53.81	
1621	493.2	463.5	54.02	
1641	499.6	469.5	53.74	
1661	525.4	493.8	53.95	
1681	496.6	466.7	53.67	
1711	508.8	478.2	53.61	
1741	500.5	470.4	53.79	
1771	510.3	479.6	53.76	
1801	513.3	482.4	53.74	
1861	535	502.8	53.75	
1981	535	502.8	53.43	
2101	536	503.8	53.23	
2221	547.7	514.8	52.99	
2341	523	491.5	53	
2461	547.9	514.9	53.06	
2581	532.3	500.3	52.78	
2701	565.8	531.8	53.4	
2801	551.8	536.1	53.28	
2813	530.2	515.1	54	
2825	548.9	533.3	55.71	
2837	753.4	731.9	57.35	
2849	959.8	932.4	58.75	
2861	1059.5	1029.3	60.11	
2873	1182.4	1148.7	60.4	
2885	1222.3	1187.5	61.01	
2897	1287.1	1250.4	61.2	
2909	1319.3	1281.7	61.3	
2921			61.5	
2933	1224	1189.1	61.4	
2945	1268.9	1232.7	61.9	71 mg/g
2957	1387.2	1347.7	62.2	
2977	1340.3	1302.1		
2997	1404.6	1364.6	63.3	
3017	1466.3	1424.5	63.4	
3037	1402.4	1362.4	64.1	
3057	1448.9	1407.6		
3077	1382	1342.6		

3097	1446.7	1405.5	64.5
3117	1478.9	1436.8	64.5
3147	1388.9	1349.3	64.8
3177	1409.9	1369.7	65
3207	1508.5	1465.5	65
3237	1488.7	1446.3	65.1
3297	1482.6	1440.3	65.26
3417	1444.1	1402.9	64.72
3537	1456.9	1415.4	65.46
3657	1544.4	1500.4	65.48
3777	1539.9	1496.0	65.53
3897	1369.1	1330.1	66.02
4017	1634.4	1587.8	65.79
4137	1497 4	1454.7	66.46

Fig 4-5, PhD Dissertation, 2/96 1500 ppmv humid feed to 48 sections of K-cloth (0.0792 g) Flow 19.8 ml/min, not insulated or heat taped

Time	GC Area	Eff. Conc.	. Ohms
0			190
10			210.8
30	3.3	2.29	249.6
50	0	0.00	252
70	0	0.00	248.6
90	0	0.00	246.4
110	0	0.00	245.5
130	0	0.00	242.8
150	0	0.00	242.2
170	0	0.00	255.4
190	0	0.00	258.9
210	0	0.00	259.8
230	0	0.00	259.9
250	0	0.00	262.1
270	0	0.00	261.9
290	4.7	3.26	263.9
310	356.4	247.50	263.3
330	1002.6	696.25	266.5
350	1614.4	1121.11	265.3
370	2041.8	1417.92	266.7
390	2077.4	1442.64	269
410	2062.1	1432.01	267
430	2212.1	1536.18	271
450	1913.3	1328.68	268.3
470	2249.6	1562.22	271.4
490	2029.4	1409.31	268.8
510	2156.4	1497.50	271.4
530	2090.5	1451.74	272.4
550	2050	1423.61	271.1
570		1563.19	272.6
590	2045.2	1420.28	270.2
610	2212	1536.11	274.6
630	2024	1405.56	271.6
650		1559.17	275.8
670		1386.60	273.2
690		1543.61	275.5
710	2066	1434.72	273

730	2265.7	1573.40	274.6
750	2091.7	1452.57	274.4
770	2218.4	1540.56	275.8
790	2136.7	1483.82	276.5
810	2221.98	1543.04	274.5
830	2145.4	1489.86	277.1
850	2112.8	1467.22	276.1
870	2151.9	1494.38	277.4
890	2065.6	1434.44	275.3
910	2032.6	1411.53	278.5
930	2168	1505.56	275.6
950	1915.8	1330.42	279
970	2127.8	1477.64	275.2
990	1864.8	1295.00	279.4
1010	2112.2	1466.81	276.3
1030	1932	1341.67	279.8
1050	2152.8	1495.00	276.7
1070	1876.9	1303.40	279.8
1090	2030.7	1410.21	277
1110	1721.2	1195.28	278.9
1130	2030.8	1410.28	278.5
1150	2220.3	1541.88	277.2
1170	2247.4	1560.69	279.3
1190	2128.7	1478.26	280.7
1210	1960.9	1361.74	280.3
1230	2145.4	1489.86	278.3

Fig 4-6, PhD Dissertation, 2/96
Resistance Changes with addition of compounds to K Cloth, Sep 95
Compounds-Dichloromethane (DCM), Methanol (MeOH), and Toluene (Tol)

		Resistance,	
Time		MeOH	MeOH
	1	72.81	1.00
	5	76.4	1.05
	7	87.3	1.20
	8	91.3	1.25
	10	98.8	1.36
	11	100.7	1.38
	12	102.33	1.41
	13	103.4	1.42
	15	106	1.46
	16	106.9	1.47
	17	107.6	1.48
	18	108.2	1.49
	20	109.5	1.50
	22	110.6	1.52
	24	111.34	1.53
	26	111.7	1.53
	28	112	1.54
	30	112.3	1.54
	32	112.6	1.55
	34	112.7	1.55
	36	112.8	1.55
	60	113.7	1.56
			Toluene
	1	76.1	0.99
	2	75.6	0.98
	4	74.2	0.96
	5	74	0.96
	6	74.8	0.97
	7	75.6	0.98
	8	76	0.98
	10	75.4	0.98
	11	74.8	0.97
	13	74	0.96
	14	77.5	1.00
	15	78.9	1.02
	16	81.2	1.05

17	80.4			1.04
19	79.7			1.03
23	86			1.11
24	84.8			1.10
				1.14
26	88.1			1.14
27	90.3			
28	92.1			1.19
29	95			1.23
30	96.5			1.25
31	103			1.33
32	109.7			1.42
33	118.7			1.54
34	122.4			1.59
35	125.7			1.63
36	126.4			1.64
37	123.3			1.60
38	123.5			1.60
48	125.4			1.62
58	123.3			1.60
			R/Ro	
			DCM	TCE
1	85.5	57.8	1.00	1.00
2	88.6	57.8	1.04	1.00
3	88	57.2	1.03	0.99
4	87.4	56.1	1.02	0.97
5	88	55.2	1.03	0.96
6	89.2	54.1	1.04	0.94
7	94.6	53	1.11	0.92
8	100.6	52.4	1.18	0.91
9	104.2	54.1	1.22	0.94
10	106.7	53.4	1.25	0.92
11	107.9	56.2	1.26	0.97
12	108.7	59.8	1.27	1.03
13	109.5	63.5	1.28	1.10
14	109.9	66.4	1.29	1.15
15	110.2	68.6	1.29	1.19
16	110.5	68.1	1.29	1.18
18	110.5	68.2	1.29	1.18
20	110.6	69	1.29	1.19
30	110.6	69.4	1.29	1.20
40	113.7	77.9	1.29	1.21
50	113.8	77.4	1.30	1.20

60	113.9	75.4	1.30	1.17
70	113.9	75.9	1.30	1.17
	113.9	75.9	1.30	1.17
			R/Ro	
			CCL4	PCE
1	63.5	80.1	1.02	1.00
2	62.2	79.9	1.00	0.99
3	61.5	80	0.99	0.99
4	61.4	79.9	0.99	0.99
5	61.8	79.6	1.00	0.99
6	64.3	79.3	1.04	0.99
7	63.2	78.5	1.02	0.98
8	60.8	78.1	0.98	0.97
9	59.1	77.8	0.95	0.97
10	62.6	78.1	1.01	0.97
11	58.3	78.1	0.94	0.97
12	64.7	78.3	1.04	0.97
13	70.4	78.3	1.13	0.97
14	68.1	79	1.10	0.98
15	67.9	79.8	1.09	0.99
16	68.5	80.6	1.10	1.00
17	69.7	80.6	1.12	1.00
18	71.5	80.8	1.15	1.00
19	71.4	81.1	1.15	1.01
20	72.4	83.4	1.17	1.04
21	74.2	86.2	1.19	1.07
22	73.9	90.2	1.19	1.12
23	73.6	90.6	1.19	1.13
24	74.1	91.9	1.19	1.14
26	74.1	96.6	1.19	1.20
27	75.1	101.1	1.21	1.26
28	74.7	106.1	1.20	1.32
30	75.1	113.3	1.21	1.41
31	76.1	115.1	1.23	1.43
32	76.3	118	1.23	1.47
34	76.8	119.2	1.24	1.48
35	77.2	121.5	1.24	1.51
36	77.4	123.7	1.25	1.54
38	77.4	124.8	1.25	1.55
39	77.5	126.6	1.25	1.57
40	77.6	124.8	1.25	1.55
41	77.7	126.2	1.25	1.57

42	77.7	126.9	1.25	1.58
43	77.7	128	1.25	1.59
45	78.3	129.6	1.26	1.61
47	78.7	134	1.27	1.66
50	78.8	136.4	1.27	1.69
52	79.1	138.1	1.27	1.72
54	79.4	139.3	1.28	1.73
56	78.4	140.1	1.26	1.74
58	78.7	144.5	1.27	1.80
60	78.8	145.2	1.27	1.80
70	79.4	149.4	1.28	1.86
80	80.8	151.2	1.30	1.88

Fig 4-7, PhD Dissertation, 2/96 Resistance/Adsorbance test results, Sep 1995 K-Cloth with DCM and TCE

Time		DCM R	TCE R	DCM R/R	TCE R/Ro		
	0	85.5	57.8	1.00	1.00		
	1	88.6	57.8	1.04	1.00		
	2	88	57.2	1.03	0.99		
	3	87.4	56.1	1.02	0.97		
	4	88	55.2	1.03	0.96		
	5	89.2	54.1	1.04	0.94		
	6	94.6	53	1.11	0.92		
	7	100.6	52.4	1.18	0.91		
	8	104.2	54.1	1.22	0.94		
	9	106.7	53.4	1.25	0.92		
	10	107.9	56.2	1.26	0.97		
	11	108.7	59.8	1.27	1.03		
	12	109.5	63.5	1.28	1.10		
	13	109.9	66.4	1.29	1.15		
	14	110.2	68.6	1.29	1.19		
	15	110.5	68.1	1.29	1.18		
	16	110.5	68.2	1.29	1.18		
	18	110.6	69	1.29	1.19		
	20	110.6	69.4	1.29	1.20		
	30	113.7	77.9	1.29	1.21		
	40	113.8	77.4	1.30	1.20		
	50	113.9	75.4	1.30	1.17		
	60	113.9	75.9	1.30	1.17		
	1					54.3	0.99
	2					54	0.99
	3					54	0.99
	4					53.65	0.98
	5					53.5	0.98
	6				,	53.3	0.98
	7					53.1	0.97
	8					53	0.97
	9					52.8	0.97
	10					52.6	0.96
	11					52.45	0.96
	12					52	0.95
	13					52	0.95
	14					51.8	0.95

15	51.5	0.94
16	51.3	0.94
17	51	0.93
19	50.65	0.93
20.5	50.3	0.92
22	50.1	0.92
23	49.85	0.91
24	49.65	0.91
26	49.3	0.90
28	48.95	0.90
30	48.75	0.89
32	48.5	0.89
34	48.2	0.88
36	48.05	0.88
38	48	0.88
40	48.3	0.88
42	48.6	0.89
44.5	49.2	0.90
47	49.45	0.91
50	49.6	0.91
53.5	50	0.92
56	50.4	0.92
57	50.7	0.93
59	51.1	0.94
60	51.3	0.94
63	51.6	0.95
66	51.9	0.95
70	52.15	0.96
82	52.4	0.96
90	52.5	0.96
100	52.5	0.96
110	52.55	0.96
120	52.6	0.96

## Water plus TCE in equal volumes in static test

Time	Ohms	
1.00	58.6	
1.67	58.5	
2.33	57.56	
3.00	56.41	
3.67	55.24	

4.33	54.39
5.00	53.95
5.67	53.91
6.33	54.09
7.00	54.34
7.67	54.2
8.33	53.97
9.00	53.87
9.67	53.71
10.33	53.75
11.00	53.67
11.67	53.67
12.33	53.7
13.00	53.86
13.67	54.2
14.33	55.04
15.00	55.05
15.67	55.06
16.33	54.27
17.00	54.41
17.67	54.47
18.33	54.8
19.00	55.09
19.67	55.01
20.33	55.5
21.00	56.54
21.67	57.24
22.33	57.66
23.00	57.64
23.67	59.48
24.33	59.47
25.00	60.22
25.67	62.37
26.33	62.18
27.00	63.21
27.67	64.46
28.33	66.44
29.00	66.9
29.67	68.32
30.33	69.95
31.00	71.02
31.67	73.04

32.33	73.95
33.00	73.13
33.67	74.64
34.33	75.48
35.00	74.7
35.67	74.5
36.33	75.14
37.00	76.69
37.67	77.62
38.33	77.63
39.00	78.12
39.67	80.85
40.33	80.71
41.00	81.31
41.67	81.44
42.33	81.09
43.00	80.13
43.67	80.46
44.33	80.34
45.00	79.77
45.67	79.9
46.33	79.93
47.00	79.02
47.67	79.08
48.33	79.24
49.00	79.54
49.67	79.42
50.33	79.25
51.00	79.69
51.67	80.23
52.33	80.13
53.00	80.68
53.67	81.05
54.33	80.05
55.00	80.35
55.67	80.86
56.33	80.13
57.00	80.59
57.67	79.99
58.33	79.54
59.00	80.03
59.67	79.54

60.33	79.16
61.00	80
61.67	79.88
62.33	79.17
63.00	80.12
63 67	80.46

# APPENDIX C

Additional Information on Thermal Catalysis of Trichloroethylene over Transition Metal Impregnated
Ambersorb 572

This appendix contains information too detailed for the professional paper in Chapter 2. The information below is summarized in Chapter 2, and the references, tables, and figures are all the same.

### AM-572-Mn

The experiments with AM-572-Mn (Table 2-6) were conducted with a dry feed at 175 °C. The catalyst appears to have an induction time as shown in Figure 2-5 where the effluent rises to about 80 percent of the influent. After the catalyst is activated, the effluent drops below detection limits (>99.9 percent conversion). During this induction time PCE was noted in the effluent, dropping off to below detection limits. Adding water to the feed reduces the TCE conversion efficiency to about 88 percent and PCE is once again to a detectable level. After 166 hours the system was turned off and the catalyst was noted to have increased in mass by about 8 percent. No chloride samples were taken to determine mineralization efficiencies.

In a separate run with AM-572-Mn with a humid feed at 210 °C and an EBCT of 3.8 s, no conversion occurred. Reducing the flow by 50 percent did not improve the performance. To determine if the temperature was too great for the oxidizing radical/carbenium ions, the temperature was reduced to 180 °C. Unfortunately this did not impact the conversion efficiency, as verified by chloride analysis. It appears that water in the system initially can impact the performance of the catalyst. The water probably blocks the access to the catalytic sites reducing or eliminating the oxidative power of this catalyst, even at elevated temperatures.

#### AM-572-Ti

The following experiments with the titanium impregnated Ambersorb 572 are summarized in Table 2-7. Using a pre-loaded AM-572-Ti at 105 °C and an EBCT of 12.7 s, no initial TCE was in the effluent. A chloride analysis showed 90 percent mineralization of TCE to HCl/Cl<sub>2</sub>. After approximately 1300 minutes the

mineralization efficiency dropped to 77 percent and reached 28 percent (conversion was down to 99.6 percent) after 1600 minutes. At this point the temperature was raised to 150 °C and flow reduced to an EBCT of 23 s. The mineralization efficiency raised up to 47 percent initially but fell to about 20 percent a short time later. Increasing the temperature again to 175 °C increased the mineralization efficiency to about 50 percent. Increasing the flow from an EBCT of 35 s to 10 s, decreased the mineralization efficiency to about 30 percent. Increasing the flow again to an EBCT of 3.7 s increased the mineralization efficiency back to approximately 50 percent indicating a possible concentration effect. The back pressure by the catalyst bed would cause a slight increase in the concentration in the bed, especially at the leading edge. This increase could, in turn, cause an increase in the catalyst performance by increasing the concentration of the reactants on the surface.

After 120 hours the catalyst was removed and had a 6 percent mass loss after the experiments. This is in direct contrast to the findings of the other low temperature runs where by-products accumulated on the surface increasing the catalyst mass. A separate aliquot of the AM-572-Ti was rinsed (up to five times) in water and the pH of the rinsate was <2 as indicated by pH paper. The catalyst/adsorbent readily wet and a tremendous amount of heat was released. After eight flushings of Milli-Q® water the rinsate finally came out at pH 7. This means that the TiCl<sub>4</sub> had not fully reacted with the water in the room air. The reaction flow may have removed the excess chlorine causing the weight loss. However, this would also put the mineralization quantities in question as to their validity.

Another portion of the original batch of AM-572-Ti was rinsed once in Milli-Q® water as described. This material, designated as AM-572-Ti(W), was dried and preloaded form for the following experiments. As the catalyst heated up to 200 °C under a TCE feed flow (EBCT of 3.3 s), approximately 240 mg TCE came off in the effluent. Subtracting the amount of TCE fed during this time and the amount destroyed (as verified by chloride analysis) this results in approximately 1/3 of the initial 100 mg TCE/g catalyst loading being desorbed. Also noted in the effluent were other by-

products. The flow was reduced (EBCT of 31 s) and the TCE effluent tailed off to non-detectable levels leaving only a small amount of PCE still in the effluent.

After a short interruption (12 hr) where the reactor cooled to room temperature without flow (and was exposed to Joule heating as part of the experiments for Chapter 3), the reactor was restarted again at 200 °C (EBCT of 11 s). The effluent continued to remain below detection limits for TCE, but now instead of PCE, small amounts of carbon tetrachloride and chloroform appeared in the effluent. After approximately 10 hours a third peak was noted, later identified as hexachloroethane (C<sub>2</sub>Cl<sub>6</sub>). A chloride analysis at this point showed the mineralization efficiency to be 58 percent. Approximately 16 hours later this had dropped to 35 percent and significant amounts of C<sub>2</sub>Cl<sub>6</sub> were in the effluent. After 30 additional hours the mineralization efficiency had dropped to 20 percent, however TCE was still non-detect in the effluent.

After 220 hours (excludes interruption) the flow was increased to obtain an EBCT of 3 s. At this flow the catalyst only lasted about 14 hours before TCE broke through and was detectable in the effluent along with a new compound (tentatively identified as pentachloroethane). However, even with TCE in the effluent and all the by-products being formed, the mineralization efficiency of the catalyst was 36 percent. The reactor was allowed to continue operating with the TCE concentration slowly increasing. As the TCE increased the C<sub>2</sub>Cl<sub>6</sub> decreased and the others remained essentially constant. A re-check of the catalytic activity a day later indicated that the mineralization efficiency had stabilized at approximately 37 percent. Total on-stream catalyst time was over 285 hours, ending with the TCE effluent concentration approximately one half of the dry 1500 ppmv feed.

#### AM-572-Co

Cobalt was the final transition metal to be used to impregnate the Ambersorb® adsorbent. The conditions reported here are summarized in Table 2-6. Initial flows

(EBCT of 7 s to 49 s) up to 145 °C showed no catalytic activity. Raising the temperature to 190 °C (EBCT of 24 s) dramatically drops the TCE effluent but also starts to show chloroform in significant quantities. A chloride analysis at the point where the TCE was approximately 20 percent of the influent as determined by the GC, indicated a mineralization efficiency of about 17 percent. The TCE concentration in

the effluent continued to drop and eventually went non-detect, with only chloroform remaining at a relatively constant level.

The flow was increased to an EBCT of 9.6 s to determine if the TCE would return to the effluent. While TCE was not seen, the chloroform levels increased slightly and slight amounts of carbon tetrachloride appeared. The mineralization efficiency went up to 47 percent. A GC/MS sample of the effluent indicated a new, barely detectable peak, phosgene. The flow was again increased to an EBCT of 2.8 s, which rapidly (within 4 hrs) yielded TCE in the effluent (4 ppmv) and also PCE in the effluent.

Reducing the reactor flow and temperature (EBCT of 32 s; 130 °C) caused the adsorption isotherms to shift, causing initial decreases in effluent concentrations. After this had re-equilibrated only chloroform, carbon tetrachloride and PCE remained in the effluent. A chloride analysis indicated a mineralization efficiency of 27 percent, even though the TCE was non-detect in the effluent. Increasing the flow to an EBCT of 9.6 s did not initially change the product spectrum, however, the mineralization efficiency decreased to 20 percent. After 2 days under these conditions TCE was noted in the effluent (2 ppmv). Another increase in flow (EBCT of 3 s) yielded significant quantities of TCE in the effluent (900 ppmv) which stayed relatively constant for the remainder of this 280 hour experiment. The catalyst mass increased by approximately 30 percent.

In a subsequent run the cobalt oxide catalyst was started at 190 °C. In this run the cobalt showed an induction period similar to the manganese as shown in Figure 2-5. The addition of water to the feed stream showed deactivation of the catalyst so the run

was stopped. No chloride samples were taken to confirm mineralization.

To determine the nature of the induction period, an aliqot of the initiall 100 g batch of AM572-Co was placed in a tube furnace and heated to 250 °C in air over night. Some of this material, AM572-Co(H), was put into the steel reactor (25 s EBCT; 280 °C). Only a 67% conversion was obtained, with chloroform and PCE as the major byproducts. The catalyst suffered a 12 percent weight loss during the run.

Since the -SO<sub>3</sub>H group can be can be ion exchanged, a new batch of catalyst was made by passing cobalt nitrate (1 M) through a bed of plain Ambersorb 572. As expected the adsorbent did not have much exchange capacity. This catalyst, AM 572-Co(e), at 180 °C (1 s EBCT) also had an induction period, after which complete conversion (>99.9 percent) is achieved, but the mineralization efficiency is only 7 percent. Shortly afterwards the catalyst completely deactivates. Heat treating the AM-572-Co(e) as above to make AM 572-Co(e)/H showed a standard adsorption curve (short loading with breakthrough) and no catalytic activity.